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Submission to the Commonwealth Department of Health Therapeutic Goods Administration

То

Amend The Scheduling of Ivermectin

Deletion of Appendix D, Item 10 from The Current S4 Poisons Scheduling

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DECLARATION:

We declare that we have no competing interests or conflicts of interest in making this submission. We represent the Australian Medical Professionals Society (AMPS). We believe the views expressed in this document are consistent with those of our members. The information provided in the submission is, as far as we know, true and accurate.

We agree to maintain confidentiality in relation to notifications of intermediate and final decisions on this consultation and submission until they are published in accordance with subsections 42ZCZP and 42ZCZS of the Therapeutic Goods Regulations Act 1990, as applicable (i.e., following referral to an expert advisory committee).

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Introduction

On 10 September 2021, a delegate of the Secretary of the Department of Health considered the advice provided by the Advisory Committee on Medicines Scheduling (ACMS) and made the decision to amend the Poisons Standard by creating a new Appendix D listing for ivermectin and thus eliminated its use as an off-label treatment option for COVID-19. This occurred with reference to subsection 52E(1) of the Therapeutic Goods Act 1989, in particular paragraph (f), which empowers the Secretary to act on any *other matters* that the Secretary considers necessary to protect public health¹. We consider this change to the Poison Scheduling for ivermectin to be inappropriate and not in the best interests of medicine in Australia².

The role of the Therapeutic Goods Administration is to apply scientific and clinical expertise to decision making, to ensure that the benefits to consumers outweigh any risks associated with the use of medicines³. However, the reasons outlined for placing constraints on the prescription of ivermectin for the treatment of COVID-19 do not appear to be based on a thorough risk benefit analysis to consumers and appear to contradict earlier authoritative safety analysis (AusPAR 2013). The use of ivermectin was restricted in a very specific context, in which the priority for public health agencies was maintaining the focus on vaccine uptake in the community, whilst maintaining control of messaging.

The Australian Medical Professionals Society (AMPS) is a growing association of medical professionals in Australia. AMPS welcomes the opportunity to make a submission to amend the scheduling of ivermectin, through deletion of Appendix D, Item 10 from the current S4 Poisons Scheduling. In seeking to provide our Society's perspective, we will discuss the set of rationales outlined by the TGA at the time of the original decision. Importantly, it is our belief that to meet our Code of Conduct obligations, we must seek to have safe, affordable and efficacious medicines available to our patients. As such, we seek to have ivermectin reinstated and available at the present time, as was the case pre-pandemic.

Prior to the amendment of September 2021, ivermectin had been available for off-label prescribing, in accordance with the clinical judgement of doctors. In a climate where clinicians became used to looking to the government for guidance on numerous pandemic-related issues in daily practice, it is true that there were no positive statements made by government bodies or associated committees, in support of the use of ivermectin for COVID-19 disease. However, many Australian doctors felt from their own analysis that the case for ivermectin was very reasonable (often in combination with other medications) and were able to use this medicine off-label, as confirmed by Minister Hunt, in a letter from August 2020⁴. Clearly no sponsor was likely to approach the TGA to seek a formal indication

¹<u>https://www.tga.gov.au/resources/publication/scheduling-decisions-final/notice-amendment-current-poisons-standard-under-paragraph-52d2a-therapeutic-goods-act-1989-0</u>

²<u>https://www.tga.gov.au/resources/publication/scheduling-decisions-final/notice-amendment-current-poisons-standard-under-paragraph-52d2a-therapeutic-goods-act-1989-0</u>

³<u>https://www.tga.gov.au/about-tga/what-we-do/role-tga#:~:text=The%20TGA%20is%20responsible%20for%20regulating%20the%20supply%2C%20import%2C%20export.be%20lawfully%20supplied%20in%20Australia.&text=The%20TGA%20is%20a%20part%20of%20the%20Australian%20Government%20Department%20of%20Health.</u>

⁴https://www.tga.gov.au/products/covid-19/covid-19-treatments/covid-19-treatments-provisional-registr ations#:~:text=Off%2Dlabel%20prescribing%20refers%20to,the%20setting%20of%20informed%20co nsent.

of ivermectin in the treatment of COVID-19 disease, given that its patent expired, but this was not a significant barrier to physician-driven off-label treatment.

With this background, the pressure and concern of the vaccine rollout and a potential negative impact on ivermectin availability, which itself implied that significant numbers of doctors were prescribing the drug, appear to have been the primary motivations in introducing Appendix D in its current form. These reasons will be discussed subsequently. As will also be discussed, AMPS members have assessed the full range of studies on ivermectin and believe that the initial hesitancy, in which claims that ivermectin was unsafe thrived, is unsupported by the overall body of literature⁵. In fact, the evidence base continues to grow that this is a safe, cost effective, efficacious and essential medicine.

In the changing context of SARS-CoV2 and COVID-19 disease, which remains prevalent despite high rates of vaccination, our view is that Australian doctors should have the maximum options available for use, based on their clinical judgement. Cognisant of our Code of Conduct obligations and placing patient care as our primary concern, we believe that ongoing restrictions on ivermectin prescribing is not suited to the current conditions of the pandemic. AMPS therefore strongly supports the deletion of Appendix D, Item 10 from the Current S4 Poisons Scheduling, in the best interest of Australian doctors and their patients.

Professional Responsibilities

AMPS has been established as a platform of advocacy for medical professionals in this country. We advocate for policies and practices which support the health and safety of the Australian public, are supremely focussed on patient care and are consistent with the Good Medical Practice Code of Conduct. The Code sets out professional obligations to ensure patient care is our highest priority. Doctors are obliged to act honestly, ethically and in a trustworthy manner. Public trust in medical professionals is a bedrock of public health. Australians expect their doctors to act competently, providing advice openly and with full disclosure and to display qualities of integrity, truthfulness, dependability and compassion⁶.

AMPS undertook a survey of membership to solicit feedback on the potential removal of Appendix D and can advise that 100% of respondents were fully supportive of the proposal to reschedule this medicine. Additionally a recent survey conducted by the Royal Australian College of General Practitioners found that the majority (54%) of doctors believe there should be no restrictions on being able to prescribe ivermectin for COVID-19⁷. Our members expressed their determination and saw it as their duty to advocate strongly for patients to have access to ivermectin, being confident of the supporting evidence-base with regard to safety, as well of its benefits in the treatment of COVID-19 disease at various stages.

In this regard, our society makes note of the 2013 AusPar Report which found no significant safety concerns reported with the use of ivermectin. Given the fiduciary obligation doctors

⁵ <u>https://ivmmeta.com/</u>

⁶file:///C:/Users/danan/Downloads/Medical-Board---Code---Good-medical-practice-a-code-of-conduct-f or-doctors-in-Australia---1-October-2020%20(15).PDF

⁷ <u>https://www1.racgp.org.au/newsgp/poll</u>

have when unwell patients present to them, to treat them to the best of their knowledge and ability, we believe that the changes to Appendix D place all doctors who are aware of the safety profile of ivermectin, in a situation which breaches our primary obligations. Furthermore, to our membership, it is of great concern that restrictions on the availability of this product has prevented vast numbers of Australians, who wished to do so, from accessing a safe treatment option that showed genuine promise.

We are not opposed to the approval and availability of other medicines for early treatment of COVID-19 disease. However, we note that decisions have been made to provisionally approved medicines with less supporting evidence than ivermectin, especially with regard to safety, and with significantly higher cost and adverse event profile, such as Remdesivir, Paxlovid and Molnupiravir⁸.

On first principles, an early treatment strategy is both separate and complementary to a vaccination strategy. However, it is now clear that mRNA vaccines have been less effective than anticipated. It is now clear that less protection is offered by currently available vaccines against new and prevailing variants of SARS-CoV2. Unfortunately, the phenomenon of waning immunity, in which protection of any kind is very limited after 4-6 months, is well documented and publicly acknowledged. With this in mind, if there was at one time a basis for a 'vaccine only strategy', it is certainly no longer the case. We believe it is now time to liberalise decision making about best clinical care to medical practitioners, who should be free to draw on their years of expertise and subject knowledge to make recommendations for the benefit of patients, at their discretion.

Given these considerations, the statement that there is not enough evidence to support the safe and effective use of ivermectin drugs (used as monotherapy or in combination with doxycycline and zinc) to prevent or treat COVID-19⁹ does not accord with the current body of evidence, amassed historically and recently. This being the case, with ivermectin being a safe and accepted item of the pharmacopoeia decades before the pandemic, we wish to highlight that the persistence of Appendix D in its current form, limits the ability of doctors to exercise their judgement on behalf of patients and thus may compromise them in their fiduciary duty to individual patients above all else.

To summarise, we have made the case that a restrictive policy regarding ivermectin does not accord with the professional opinions of our membership, nor with a large proportion in the wider medical community. We believe that, in practice, such a policy contradicts our Codes of Conduct and wish to highlight that this can be remedied by the deletion of Appendix D, Item 10.

⁸https://mail-attachment.googleusercontent.com/attachment/u/0/?ui=2&ik=614ed1668c&attid=0.1&permmsgid=msg-f:1743540180858574027&th=18324c8e8bb214cb&vi ew=att&disp=inline&realattid=f_17v5qnoy0&sadnir=2&saddbat=ANGjdJ9BgyYZBQpBGpL4ZmYvV9fj1xekMYbqPr_qyoMkcsvEmyvwaYSrl1qXRmi82_s0BUm-cvwQkYMU EKh0Fm7qoGs8ZhqqmZOYG1YR6_N0K36uxNFu59R3E5PzUOniupo130hjZJqUoY8MANRcBcVhEWqCXZVu_mZSPM2QPOYPL89E3sicJn5bnYZIFY6sShMz-yFl8d58 h0qdsK_WQ8srP8JyvoLtoQg5leIAc2D4DukO2P_t8BpqII5vA-WTdx027GC555t7fxgq9ybld74vazzE8RVSzwIprirguuK2qtEVR0-V934pvQ0oSVchpkWahuJ5KI=hcNym62Ty -0dci4KBvHipT8SWbq-M8hMq0JvYLqVhIBkncnZZzp9NDmg9-BINV-CawttzoC8tSylaKrKFbBnlukBYd8csp638a_c1u3sGW8cSpcjex40J=eCHM7W3m1jope=6f4P2lcPL=8K d0OhXkg-kUxPLZ2Vpaoom-zYLxWPbfAX80B8bhffWtWsoZ9Rii1li5o-tJSjYpBPp1CbOjWoemCGHJIQTEvQleOF4XfKQiPpvMZwFY7iJ2wVJ4mgKEsYxHiZNSysC5hSOX URdX1k44AFUK8sDdry6mcdcsYPysHagyxe9gRrA17eNkc1Jtn34qkOh1YtWw4Ocmx9Lj8iRL=8QX70S-0eh5TaHQfls_bltcuQtaVB_uU1SzAnHOwGvAJr1Qd7sSRRIPuCyv vsjd8eMzYVQW0bW2P0iEhcnPyKY

⁹ <u>https://www.health.gov.au/health-alerts/covid-19/treatments/about</u>

Reasons given for the Rescheduling of Ivermectin

On 10 September 2021 a delegate of the Secretary of the Department of Health considered the advice provided by the Advisory Committee on Medicines Scheduling (ACMS) and decided to amend the Poisons Standard by creating a new Appendix D listing for ivermectin, in effect banning it for use as an off-label treatment option for COVID-19. In statements made by the TGA¹⁰, this change to Poison Scheduling was backed up with reference to subsection 52E(1) of the Therapeutic Goods Act 1989, paragraph (f)¹¹, together with 3 stated reasons relating to public health, considered in the remainder of this section:

Reason 1. Serious concerns that there are significant public health risks associated with the prescribing of ivermectin for COVID-19. This includes the likelihood that people who have been prescribed the substance for this purpose may believe themselves to be protected from the disease and not get vaccinated or tested and seek appropriate medical care if they develop symptoms.

Reason 2. Potential to cause severe adverse events in persons, particularly when taken in high doses that have recently been described in social media and other sources for the prevention or treatment of COVID-19 infection.

Reason 3. Concern that if action is not taken to address these concerns, it is possible that oral ivermectin will be in shortage in Australia for the treatment of the conditions for which it has been properly evaluated and approved in accordance with scientific data.

AMPS does not believe Reason 1 justified the prohibition of ivermectin prescribing for the treatment of COVID-19. We believe that every intervention has to be judged on its own merits and that doctors and patients should be able to make these decisions together, in an atmosphere free from undue pressure for any other party. We further believe that the decision of an individual to be vaccinated is a separate and complementary one to any treatment strategy employing ivermectin. Regarding Reason 3, supply has not been reported to be a problem in Australia or world-wide.

AMPS is of the understanding that the role of the TGA is to determine the safety of medicines and regulate products based on an assessment of risks against benefits¹²¹³. In this spirit, we do not take the view that the legislative provisions within the Therapeutic Goods Act necessarily allow the TGA to restrict access to acceptable pre-existing medical options, as a means of encouraging public behaviour to meet other policy objectives. This

¹⁰<u>https://www.tga.gov.au/resources/publication/scheduling-decisions-final/notice-amendment-current-poisons-standard-under-paragraph-52d2a-therapeutic-goods-act-1989-0</u>

¹¹https://www.tga.gov.au/resources/publication/scheduling-decisions-final/notice-amendment-current-p oisons-standard-under-paragraph-52d2a-therapeutic-goods-act-1989-0

¹²https://www.tga.gov.au/about-tga/what-we-do/role-tga#:~:text=The%20TGA%20is%20responsible% 20for%20regulating%20the%20supply%2C%20import%2C%20export,be%20lawfully%20supplied%2 0in%20Australia.&text=The%20TGA%20is%20a%20part%20of%20the%20Australian%20Governmen t%20Department%20of%20Health.

¹³<u>https://www.tga.gov.au/how-we-regulate/advertising/legal-framework/act-regulations-and-code-offen</u> ces/how-tga-regulates

kind of justification, implicitly present in Reason 1 with regard to uptake of provisionally approved vaccines, was not subjected to wide consultation.

As stated previously, early treatment is a separate and complementary strategy, which can and does coexist in the treatment of Australians facing COVID-19 disease. In regards to currently available provisionally approved vaccines against SARS-Cov2, we note with significant concern, the unprecedented rates of adverse event reports, including deaths, injury and disablement, being seen in Australia and across the world¹⁴. Regardless of this, however, we note that consequent to Appendix D, Item 10, vaccinated Australians who suffer COVID-19 are currently being denied access to the full range of early treatment options, despite the objective of high vaccination rates having already been achieved in Australia.

Also with regarding Reason 2, in consideration of safety, the (NCCET) conducted a review of the clinical data regarding the use of ivermectin in the management of COVID-19 and concluded;

"The available research evidence does not yet provide reasonable certainty to recommend for or against the use of ivermectin and therefore the Taskforce recommends ivermectin not be used outside of randomised trials. The certainty of the current evidence base varies from low to very low.¹⁵"

We note that the term "reasonable certainly" is ambiguous in terms of drug regulation, in that the threshold for what is reasonable is not defined and may be viewed differently by different parties. We point out that this recommendation has been challenged by experts both National and Internationally¹⁶¹⁷¹⁸. We believe there is ample controlled evidence to support the effectiveness of ivermectin both alone and in combination, in addition to the notable documented experience of countries such as India and Peru, in which a strong correlation has been reported between ivermectin use and mortality reductions. Nevertheless, efficacy was not a reason outlined by the TGA as a consideration in the scheduling decision¹⁹²⁰²¹.

Therefore, in the following section, we will focus on addressing safety concerns and the claim that ivermectin has the potential to cause severe adverse events, pertinent to Reason

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https://8630368.fs1.hubspotusercontent-na1.net/hubfs/8630368/AMPS/Altman%20Report%20Final%20Ve rsion%2011-8-22%20(1).pdf?utm_source=hs_email&utm_medium=email&_hsenc=p2ANqtz-8HS0cEyUJu QHjoxCYMYvaYAqn1CWxMNk_F4VyGSiymi6QxgE6AEh9SJNXh6yR0hIVEAxxC

15 https://covid19evidence.net.au/wp-content/uploads/NC19CET_Published_Guideline_V48_0.pdf

20 https://osf.io/9egh4/

ep-deaths-low-7311786/

¹⁶file:///C:/Users/danan/Downloads/COCHRANE%20Fordham-Review%20of%20Cochrane%20Report%20copy.pdf

¹⁷ <u>https://quadrant.org.au/opinion/qed/2021/08/commentary-on-nccet-statement-on-ivermectin/</u>

¹⁸ https://quadrant.org.au/opinion/public-health/2021/10/we-cant-vaccinate-this-pandemic-away/ 19 https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8383101/

²¹https://indianexpress.com/article/cities/lucknow/uttar-pradesh-government-says-ivermectin-helped-to-ke

2, above. We will focus on peer reviewed data, rather than the potential dangers associated with data sourced from social media posts, as considered in the TGA reasoning.

Ivermectin Safety and Clinical Benefits

The essential issue is that in the case of a repurposed compound with documented safety and excellent tolerability, such as ivermectin, doctors should not be hindered in evaluating such pre-existing treatments and adopting them if they so choose, in pursuit of the best care of their patients. This simply reflects a reasonable and time-honoured approach, employing critical appraisal, risk benefit analysis and informed consent, in keeping with good medical practice. In the context of a novel health concern, we argue that a responsible physician-directed process is eminently suitable when the compounds under consideration are familiar to doctors and have excellent known safety profiles. This is the case with ivermectin, especially where considered in the pre-hospital phase of COVID-19 treatment, where other options have been more limited.

As outlined in the TGA's 2013 AusPar Report for ivermectin, no significant safety concerns were found with the use of ivermectin²². Very importantly, the report found no safety concerns even at 10 times the (then) current approved dose of 200ug/kg²³. The U.S. National Institute of Health (NIH) has recognised that "ivermectin has been widely used and is generally well tolerated"²⁴. A recent systematic review stated "ivermectin at the usual doses is considered extremely safe for use in humans"²⁵. In 2018, ivermectin was added to the WHO list of Essential Medicines and in supporting the submission for inclusion in the list, the WHO concluded that the adverse events associated with ivermectin are "*primarily minor and transient*". The clinical evaluator in the WHO Report found that there were no significant safety concerns or serious adverse events reported with the use of ivermectin²⁶.

In February 2021, an expert toxicology report on the safety of ivermectin was collated based on a review of over 500 articles. This unprecedented work is well worth considering in detail and outlined the following:

"Hundreds of millions of human subjects have been treated with ivermectin for curative or prophylactic purposes worldwide over the last 3 decades. The reference list of this report demonstrates that a large body of data is available, which allows for a detailed analysis of ivermectin medical safety....

https://www.covid19treatmentguidelines.nih.gov/therapies/antiviral-therapy/ivermectin/

https://journals.lww.com/americantherapeutics/fulltext/2021/08000/ivermectin

²² https://www.tga.gov.au/sites/default/files/auspar-ivermectin-131030.pdf

²³ Australian Public Assessment Report for Ivermectin – October 2013 <u>https://www.tga.gov.au/auspar/auspar-ivermectin</u>

²⁴ National Institutes of Health, COVID-19 Treatment Guidelines: ivermectin,

²⁵ Andrew Bryant et al., Ivermectin for Prevention and Treatment of COVID-19 Infection: A Systematic Review, Meta-analysis, and Trial Sequential Analysis to Inform Clinical Guidelines, 28 American Journal of Therapeutics 434, 435 (Jul./Aug. 2021), available at

²⁶ WHO Expert Committee on the Selection and Use of Essential Medicines: Application for inclusion of ivermectin on the WHO Model List of Essential Medicines (EML) and Model List of Essential Medicines for Children (EMLc) for the indication of Scabies at 19 (Dec. 2018)

...Taking into account all the above, the author of the present analysis of the available medical data concludes that the safety profile of ivermectin has so far been excellent in the majority of treated human patients so that ivermectin human toxicity cannot be claimed to be a serious cause for concern²⁷."

In this regard, a decisive legal opinion from the U.S. Nebraska State Attorney General's Office (14 October 2021) is highly instructive. It provided a detailed analysis of the arguments regarding ivermectin and off-label prescribing and a copy of this ruling forms Annexure 1 to this Submission. The Co-signatories rely upon this opinion in full as it pertains to ivermectin.

The opinion states in part:

"The data show not only that the adverse side effects are minor, but also that the percentage of people who report experiencing any adverse events is vanishingly small. The latest statistics available through VigiAccess report only 5,674 adverse drug reactions from ivermectin between 1992 and October 13, 2021. This number is incredibly low considering that "more than 3.7 billion doses" of ivermectin have been administered to humans worldwide since the 1980s²⁸."

The brief but comprehensive review of the safety of ivermectin provided here does not provide any clear or convincing evidence that ivermectin poses such a threat to public health and safety that it required sudden rescheduling in the middle of a pandemic as a poison when prescribed for COVID-19. In truth, no data exists in support of serious harm. It is likely that the absence of safety concerns relating to ivermectin was the very reason for the rapid commencement of multiple early controlled trials in COVID-19 disease overseas, after widespread interest in the potential benefits of this highly versatile drug.

Ivermectin has documented pharmacological mechanisms that led clinicians to believe this extremely safe medicine could be repurposed effectively for the treatment of COVID-19. It has been known for over 10 years that ivermectin demonstrated antiviral activity against several RNA viruses by blocking the nuclear trafficking of viral proteins²⁹. A comprehensive systematic review summarises the antiviral effects of ivermectin, including in vitro and in vivo studies over the past 50 years ³⁰. Another paper titled, "Ivermectin: an award-winning drug with expected antiviral activity against COVID-19" put forward that Ivermectin, an FDA-approved broad-spectrum antiparasitic agent, had demonstrated antiviral activity against a number of DNA and RNA viruses, including severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2)³¹. As well as ivermectin's antiviral benefits there is also research literature that outlines its recognised "anti-inflammatory capacity"³².

²⁷ Descotes, J. Expert Review Report – Medical Safety of Ivermectin. 3 March 2021 https://www.medincell.com/wp-content/uploads/2021/03/Clinical_Safety_of_Ivermectin March_2021.pdf

²⁸ U.S. Nebraska State Attorney General opinion. Prescription of Ivermectin or hydroxychloroquine as Off-Label medicines for the Prevention or Treatment of Covid-19. 14 October 2021

²⁹ https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7539925/

³⁰ <u>https://pubmed.ncbi.nlm.nih.gov/32533071/</u>

³¹ https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7539925/

³² https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7476419/

A review titled "Emerging Evidence Demonstrating the Efficacy of Ivermectin in the Prophylaxis and Treatment of COVID-19" concluded:

"Meta-analyses based on 18 randomized controlled treatment trials of ivermectin in COVID-19 have found large, statistically significant reductions in mortality, time to clinical recovery, and time to viral clearance. Furthermore, results from numerous controlled prophylaxis trials report significantly reduced risks of contracting COVID-19 with the regular use of ivermectin. Finally, the many examples of ivermectin distribution campaigns leading to rapid population-wide decreases in morbidity and mortality indicate that an oral agent effective in all phases of COVID-19 has been identified ³³."

Finally, an online real time meta-analysis of the clinical safety and efficacy of ivermectin in COVID-19 disease is well worth considering and can be found at <u>www.ivmmeta.com</u>: as of 9 September 2022, this includes 91 studies, of which 41 were randomised controlled trials involving 11,141 patients³⁴). This resource illustrates the high level of international interest in the clinical submission of ivermectin for potential use in COVID-19. When taken in totality, the clinical data presented at <u>www.ivmmeta.com</u> presents a compelling case for the safety and efficacy of ivermectin. More than 20 countries (including India, Mexico, regions of Peru, Argentina, Japan, Dominican Republic and Brazil) have adopted ivermectin for the management of COVID-19. Collectively, the studies strongly suggest that "ivermectin reduces the risk for COVID-19 with very high confidence for mortality, ventilation, ICU admission, hospitalisation, progression, recovery, [number of] cases, viral clearance, and in pooled analysis... Meta-analysis using the most serious outcome measure shows 62% [57-70%] and 83% [74-89%] improvement for early treatment and prophylaxis".

At this stage, public health officials and the medical profession generally have had time to review the accumulating data regarding ivermectin, in addition to the rapid mutation rate of the SARS-CoV2 and waning vaccine efficacy. We believe it is vital to reconsider the role of ivermectin in the arsenal of available drugs. It is important to point out that we are not aware of any other occasion on which an established drug in the Australian pharmacopoeia that has previously been considered very safe, has been rescheduled in such a way as to make its prescription illegal for doctors.

AMPS can find no clear and conclusive evidence to support the TGA claims that ivermectin poses a safety risk to the public with the potential for a high incidence of severe adverse events. Rather, our review of the evidence demonstrates that ivermectin is a fully approved, AurPar reviewed, Nobel prize winning WHO essential medicine, that has been given in billions of doses with minimal adverse reaction reported. We consider that Australian doctors should again be afforded professional discretion with regard to ivermectin use, which may translate to benefit in future seasonal outbreaks of SARS-CoV2/COVID-19 disease, with flow-on benefits to the hospital system, with very little downside, as we have summarised.

³³<u>https://journals.lww.com/americantherapeutics/fulltext/2021/06000/review_of_the_emerging_evidenc</u> <u>e_demonstrating_the.4.aspx</u>

³⁴ https://ivmmeta.com/

Conclusion

AMPS believes in the primacy of the doctor/patient relationship within medicine and stands firmly opposed to the placement of excessive constraint on the clinical judgement of doctors. Now that Australian vaccination rates have risen to such high levels, we assert that it is consistent at this time to freshly reevaluate historic decisions in the full light of today's context.

In making this submission, foremost in the thinking of our Society is that ivermectin cannot be construed to be a hazard to the health of the Australian people. This assertion contradicts the most extensive drug safety review of ivermectin in the literature³⁵, the well known evaluation of the WHO in 2018³⁶, the decisive legal opinion of the Nebraska State Attorney General's Office³⁷, as well as the TGA's own 2013 AusPar Report³⁸. As such, we contend that in the current context, the use of off-label ivermectin cannot plausibly be said to constitute a threat to the public health of Australians, in the spirit of subsection 52E(1) of the Therapeutic Goods Act 1989, particular paragraph (f)³⁹.

As a Society, we applaud this move of the TGA to open consultation with regard to Appendix D, Item 10. AMPS believes that the continuing restriction of ivermectin would at this stage represent a serious error in judgement. In this regard, we draw attention to the humility recently expressed by Dr Rochell Walensky the Director of the CDC told, who told employees recently:

*"To be frank, we are responsible for some pretty dramatic, pretty public mistakes from testing, to data, to communications*⁴⁰*"*

As we have outlined in this document, we consider that the Australian Regulators now have the opportunity to reconsider these questions, in a way which is not only likely to benefit the health of Australians, but reinforce the invaluable role of doctors' clinical judgement and expertise in the use of safe repurposed therapies in individualised patient care.

³⁵ Descotes, J. Expert Review Report – Medical Safety of Ivermectin. 3 March 2021 https://www.medincell.com/wp-content/uploads/2021/03/Clinical_Safety_of_Ivermectin March_2021.pdf

³⁶ WHO Expert Committee on the Selection and Use of Essential Medicines: Application for inclusion of ivermectin on the WHO Model List of Essential Medicines (EML) and Model List of Essential Medicines for Children (EMLc) for the indication of Scabies at 19 (Dec. 2018)

³⁷ U.S. Nebraska State Attorney General opinion. Prescription of Ivermectin or hydroxychloroquine as Off-Label medicines for the Prevention or Treatment of Covid-19. 14 October 2021 38 https://www.tga.gov.au/sites/default/files/auspar-ivermectin-131030.pdf

³⁹<u>https://www.tga.gov.au/resources/publication/scheduling-decisions-final/notice-amendment-current-poisons-standard-under-paragraph-52d2a-therapeutic-goods-act-1989-0</u>

⁴⁰ https://www.washingtonpost.com/opinions/2022/08/18/cdc-changes-next-pandemic-preparation/

Annexure-1



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DOUGLAS J. PETERSON ATTORNEY GENERAL

TATE OF NEBRASK OFFICIAL OCT 1 4 2021 DEPT. OF JUSTICE

SUBJECT: Prescription of Ivermectin or Hydroxychloroquine as Off-Label Medicines for the Prevention or Treatment of Covid-19

- REQUESTED BY: Dannette R. Smith Chief Executive Officer Nebraska Department of Health and Human Services
- WRITTEN BY: Douglas J. Peterson, Attorney General James A. Campbell, Solicitor General Mindy L. Lester, Assistant Attorney General

INTRODUCTION

On September 16, 2021, you requested our opinion on whether it would be "deemed unlawful or otherwise subject to discipline under [Neb. Rev. Stat. § 38-186] for an appropriately licensed health care provider, once informed patient consent has been appropriately obtained, to prescribe" ivermectin, hydroxychloroquine, or other "off label use" medications "for the treatment or prevention of COVID-19." You requested this opinion in your role as Chief Executive Officer of the Nebraska Department of Health and Human Services ("Department"). Neb. Rev. Stat. § 84-205(4) gives you, as the head of an executive department, the authority to ask our office's opinion on legal questions like this one.

The Department, acting through its Division of Public Health, enforces the Nebraska Uniform Credentialing Act ("UCA"). The purpose of the UCA is to protect public

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health, safety, and welfare.¹ One way in which the Department protects the public is by investigating complaints alleging that licensed healthcare professionals have committed UCA violations.² After the Department completes an investigation, it refers the matter to the appropriate professional board to consider and make a recommendation to the Attorney General. Neb. Rev. Stat. § 38-186 then gives the Attorney General the authority to file a petition for discipline against the healthcare provider if such action is warranted.

You indicate in your request that "[c]onsumers and health care providers have been and continue to be inundated with information and opinions[] regarding COVID-19 treatment and prevention." You also note that due to the "sheer volume" of conflicting information, questions have been raised "regarding the permissibility of certain medications for the treatment or prevention of COVID-19." This observation is consistent with questions that our office has received from constituents and discussions that our office has witnessed at some of the professional boards' meetings.

After receiving your question and conducting our investigation, we have found significant controversy and suspect information about potential COVID-19 treatments. A striking example features one of the world's most prestigious medical journals—the Lancet. In the middle of the COVID-19 pandemic, the Lancet published a paper denouncing hydroxychloroquine as dangerous.³ Yet the reported statistics were so flawed that journalists and outside researchers immediately began raising concerns.⁴ Then after one of the authors refused to provide the analyzed data, the paper was retracted,⁵ but not before many countries stopped using hydroxychloroquine and trials were cancelled or interrupted. The Lancet's own editor in chief admitted that the paper was a "fabrication," "a monumental fraud,"⁶ and "a shocking example of research misconduct in the middle of

⁴ Melissa Davey, Questions raised over hydroxychloroquine study which caused WHO to halt trials for Covid-19, The Guardian (May 27, 2020), available at <u>https://www.theguardian.com/science/2020/may/</u> <u>28/questions-raised-over-hydroxychloroquine-study-which-caused-who-to-halt-trials-for-covid-19</u> (last visited Oct. 14, 2021).

⁵ Sarah Boseley & Melissa Davey, Covid-19: Lancet retracts paper that halted hydroxychloroquine trials, The Guardian (Jun. 4, 2020), available at <u>https://www.theguardian.com/world/2020/jun/04/covid-19lancet-retracts-paper-that-halted-hydroxychloroquine-trials</u> (last visited Oct. 14, 2021).

⁶ Roni Caryn Rabin, The Pandemic Claims New Victims: Prestigious Medical Journals, New York Times (Jun. 14, 2020), available at <u>https://www.nytimes.com/2020/06/14/health/virus-journals.html</u> (last visited Oct. 14, 2021).

¹ Neb. Rev. Stat. § 38-128(1).

² Neb. Rev. Stat. § 38-1,124.

³ Mandeep R. Mehra et al., Hydroxychloroquine or chloroquine with or without a macrolide for treatment of COVID-19: a multinational registry analysis, The Lancet (May 22, 2020), available at <u>https://www.thelancet.com/action/showPdf?pii=S0140-6736%2820%2931180-6</u> (last visited Oct. 14, 2021).

a global health emergency."⁷ When fraudulent information is published in a leading medical journal, it understandably leads to skepticism in some physicians and members of the public. Mindful of these concerns about misunderstandings and mistrust, we have drafted a rather lengthy opinion that aims to address the public confusion and outline the relevant scientific literature that supports our legal conclusions.

At the outset, we pause to delineate the parameters of this opinion. The question presented asked about ivermectin, hydroxychloroquine, and other drugs used "off label"that is, for a purpose other than the specific use approved by the U.S. Food and Drug Administration ("FDA"). To enable us to respond in a timely manner, we have confined our discussion to ivermectin and hydroxychloroquine only. But in doing so, we do not mean to rule out the possibility that other off-label drugs might show promise-either now or in the future—as a prophylaxis or treatment against COVID-19. Also, because our investigation has revealed that physicians who currently use hydroxychloroquine for COVID-19 do so as either a prophylaxis or an early treatment for outpatients (as opposed to a late treatment in hospitalized patients), we will confine our consideration of hydroxychloroquine to those two uses. In addition, we note that there are treatment options the FDA has approved, either through an Emergency Use Authorization ("EUA") or through the regular FDA drug-approval process, for COVID-19 prophylaxis or treatment. These include monoclonal antibodies, vaccines, and remdesivir. We do not take any position on those options because they are outside the scope of the question asked.

In the end, as we explain below, we find that the available data does not justify filing disciplinary actions against physicians simply because they prescribe ivermectin or hydroxychloroquine to prevent or treat COVID-19. If, on the other hand, healthcare providers neglect to obtain informed consent, deceive their patients, prescribe excessively high doses, fail to check for contraindications, or engage in other misconduct, they might be subject to discipline. But based on the evidence that currently exists, the mere fact of prescribing ivermectin or hydroxychloroquine for COVID-19 will not result in our office filing disciplinary actions. While our terminology throughout this opinion focuses on physicians prescribing these medicines, what we conclude necessarily applies to other licensed healthcare professionals who prescribe, participate in, or otherwise assist with a treatment plan utilizing these medications.

ANALYSIS

1. The Nebraska Uniform Credentialing Act and Other Relevant Law

The UCA was enacted by the legislature to license and regulate persons and businesses that provide healthcare and health-related services.⁸ The UCA was adopted

⁷ Boseley & Davey, supra.

⁸ Neb. Rev. Stat. §§ 38-102 & 38-104.

to protect public health, safety, and welfare, and to provide for the efficient, adequate, and safe practice of credentialed persons and businesses.⁹ "It is the intent of the Legislature," the UCA explains, "that quality health care services and human services be provided to the public" and "that professionals be regulated by the state only when it is demonstrated that such regulation is in the best interest of the public."¹⁰

The UCA grants the Director of Public Health of the Department's Division of Public Health the authority to deny a credential, refuse a credential renewal, or discipline a credential holder, although the Chief Medical Officer (if one is appointed) shall perform the Director's duties for decisions in contested administrative cases.¹¹ The Department must provide "the Attorney General with a copy of all complaints it receives and advise the Attorney General of investigations it makes" regarding possible violations of the UCA.¹² Following review and recommendation from the appropriate professional health board, the Attorney General must then determine whether the credential holder has violated any statutes or regulations and decide whether to proceed with administrative action.¹³

If the Attorney General determines that a violation has occurred, he "shall" file a petition for disciplinary action with the Department.¹⁴ The Attorney General cannot prevail in disciplinary proceedings against a licensed healthcare professional unless he proves the claim by clear and convincing evidence.¹⁵

The grounds for disciplinary action are set forth in Neb. Rev. Stat. § 38-178 and include, among other things, acting with "gross incompetence or gross negligence," practicing in "a pattern of incompetent or negligent conduct," or engaging in "unprofessional conduct" as set forth in Neb. Rev. Stat. § 38-179.¹⁶ Gross incompetence is a very high standard; it occurs only when there is "such an extreme deficiency on the part of a physician in the basic knowledge and skill necessary for diagnosis and treatment that one may reasonably question his or her ability to practice medicine at the threshold level of

- 11 Neb. Rev. Stat. §§ 38-176(1) & 38-1,101.
- ¹² Neb. Rev. Stat. § 38-1,107(1).
- ¹³ Neb. Rev. Stat. §§ 38-1,107 & 38-1,108.
- ¹⁴ Neb. Rev. Stat. § 38-186.

¹⁵ Poor v. State, 266 Neb. 183, 190, 663 N.W.2d 109, 115 (2003); Davis v. Wright, 243 Neb. 931, 936-37, 503 N.W.2d 814, 818 (1993).

¹⁶ Neb. Rev. Stat. § 38-178(6), (24).

⁹ Neb. Rev. Stat. § 38-103.

¹⁰ Neb. Rev. Stat. § 38-128(1).

professional competence."¹⁷ Neb. Rev. Stat. § 38-179 generally defines unprofessional conduct as a "departure from or failure to conform to the standards of acceptable and prevailing practice of a profession or the ethics of the profession, regardless of whether a person, consumer, or entity is injured, or conduct that is likely to deceive or defraud the public or is detrimental to the public interest."¹⁸ Along these same lines, the regulation governing physicians states that unprofessional conduct includes:

[c]onduct or practice outside the normal standard of care in the State of Nebraska which is or might be harmful or dangerous to the health of the patient or the public, not to include a single act of ordinary negligence.¹⁹

Healthcare providers do not violate the standard of care when they "select between two reasonable approaches to . . . medicine."²⁰ Regulations also indicate that physicians may utilize reasonable "investigative or unproven therapies" that reflect a reasonable approach to medicine so long as physicians obtain "written informed patient consent."²¹ "Informed consent concerns a doctor's duty to inform his or her patient," and it includes telling patients about "the nature of the pertinent ailment or condition, the risks of the proposed treatment or procedure, and the risks of any alternative methods of treatment, including the risks of failing to undergo any treatment at all."²² Regulations require physicians "to keep and maintain" records that disclose the "advice and cautionary warnings provided to the patient."²³

Prescribing medicines for off-label use—that is, for some purpose other than the use approved by the FDA—often falls within the standard of care. Indeed, "[o]ff-label use is legal, common, and necessary,"²⁴ and "[c]ourts have repeatedly recognized the propriety of off-label use."²⁵ This includes the U.S. Court of Appeals for the Eighth Circuit, which has acknowledged that "[d]octors may prescribe an FDA-approved drug for

¹⁷ Langvardt v. Horton, 254 Neb. 878, 895, 581 N.W.2d 60, 70-71 (1998).

¹⁸ Neb. Rev. Stat. § 38-179.

¹⁹ 172 Neb. Admin. Code § 88-009(Q).

²⁰ Whittle v. Dep't of Health & Hum. Servs., 309 Neb. 695, 721-22, 962 N.W.2d 339, 356-57 (2021).

²¹ 172 Neb. Admin. Code § 88-009(B).

²² Curran v. Buser, 271 Neb. 332, 337, 711 N.W.2d 562, 568 (2006) (citations omitted).

²³ 172 Neb. Admin. Code § 88-009(B).

²⁴ James M. Beck & Elizabeth D. Azari, FDA, Off-Label Use, and Informed Consent: Debunking Myths and Misconceptions, 53 Food & Drug L.J. 71, 76 (1998) (capitalization omitted).

²⁵ Id. (collecting cases).

nonapproved uses."²⁶ And the U.S. Supreme Court, in an analogous context, has affirmed that "off-label' usage of medical devices" is an "accepted and necessary" practice.²⁷ Even the FDA recognizes that off-label use is legitimate: it has said for many decades that once it approves a drug, "a physician may prescribe it for uses or in treatment regimens or patient populations that are not included in approved labeling."²⁸ Expanding on that point, the FDA has explained that "healthcare providers generally may prescribe [a] drug for an unapproved use when they judge that it is medically appropriate for their patient."²⁹ Nothing in the federal Food, Drug, and Cosmetic Act ("FDCA") "limit[s] the manner in which a physician may use an approved drug."³⁰

Based on these principles, we conclude that governing law allows physicians to use FDA-approved medicines that are unproven for a particular off-label use so long as (1) reasonable medical evidence supports that use and (2) a patient's written informed consent is obtained. In the context of this ever-changing global pandemic, we note that it is appropriate to consider medical evidence outside of Nebraska and to give physicians who obtain informed consent an added measure of deference on their assessment of the available medical evidence.

2. COVID-19 and SARS-CoV-2

The disease known as COVID-19 and the virus that causes it—SARS-CoV-2 took the world by storm in late 2019 and early 2020. While there is still so much that the medical community does not know about SARS-CoV-2 and COVID-19, it is widely recognized that COVID-19 is a multifaceted disease. "[A]dults with SARS-CoV-2 infection can be grouped" into at least three different categories depending on the progression of their disease.³¹ The first group has an asymptomatic or presymptomatic infection, meaning that those individuals have "test[ed] positive for SARS-CoV-2" but "have no symptoms

27 Buckman Co. v. Plaintiffs' Legal Comm., 531 U.S. 341, 350 (2001).

²⁸ FDA Drug Bulletin at 5 (Apr. 1982), available at <u>https://play.google.com/books/reader?</u> id=3f3YC3Gw6sEC&pg=GBS.PA6&hl=en (last visited Oct. 14, 2021).

²⁶ Rhone-Poulenc Rorer Pharms., Inc. v. Marion Merrell Dow, Inc., 93 F.3d 511, 514 n.3 (8th Cir. 1996).

²⁹ U.S. Food & Drug Administration, Understanding Unapproved Use of Approved Drugs "Off Label" (Feb. 5, 2018), <u>https://www.fda.gov/patients/learn-about-expanded-access-and-other-treatment-options/understanding-unapproved-use-approved-drugs-label</u> (last visited Oct. 14, 2021).

³⁰ FDA Drug Bulletin, *supra*, at 5. Because the question posed to us asks about prescribing drugs for off-label use, any view on the legality of efforts to market drugs for off-label use is outside the scope of this opinion.

³¹ National Institutes of Health, Clinical Spectrum of SARS-CoV-2 Infection, COVID-19 Treatment Guidelines (Apr. 21, 2021), available at <u>https://www.covid19treatmentguidelines.nih.gov/overview/clinical-spectrum/</u> (last visited Oct. 14, 2021).

that are consistent with COVID-19."³² A second group experiences a mild illness that manifests itself through "any of the various signs and symptoms of COVID-19 (e.g., fever, cough, sore throat, malaise, headache, muscle pain, nausea, vomiting, diarrhea, loss of taste and smell)" but does not include "shortness of breath, dyspnea, or abnormal chest imaging."³³ And a third group suffers from a more severe illness marked by "evidence of lower respiratory disease" and deficient "oxygen saturation" levels.³⁴ When people in this third category reach a critical level, they often "have respiratory failure, septic shock, and/or multiple organ dysfunction."³⁵

A recently published paper on COVID-19 recognized that "for reasons that are yet to be clarified, early treatment has not been emphasized" in Western countries like the United States.³⁶ Despite this, many healthcare providers in the United States advocate for early treatment, particularly for high-risk patients. In fact, scores of treating and academic physicians have published papers in well-respected journals like the American Journal of Medicine explaining that the "multifaceted pathophysiology of life-threatening COVID-19 illness . . . warrants early interventions"³⁷ and encouraging "outpatient treatment of the illness with the aim of preventing hospitalization or death."³⁸ Also, a declaration of the International Alliance of Physicians and Medical Scientists—which is apparently signed by over 10,000 physicians and scientists, more than 60 of whom are publicly identified online—supports a doctor's choice to provide early COVID-19 care rather than "advising their patients to simply go home . . . and return when their disease worsens."³⁹

32 Id.

33 Id.

34 Id.

35 Id.

Matthieu Million et al., Early combination therapy with hydroxychloroquine and azithromycin reduces mortality in 10,429 COVID-19 outpatients, 22 Reviews in Cardiovascular Medicine 1063, 1063 (Sept. 2021), <u>https://rcm.imrpress.com/article/2021/2153-8174/2153-8174-22-3-1063.shtml</u> (last visited Oct. 14, 2021).

³⁷ Peter A. McCullough et al., Multifaceted highly targeted sequential multidrug treatment of early ambulatory high-risk SARS-CoV-2 infection (COVID-19), 21 Reviews in Cardiovascular Medicine 517, 518 (Dec. 2020), available at <u>https://rcm.imrpress.com/article/2020/2153-8174/RCM2020264.shtml</u> (last visited Oct. 14, 2021) (including 57 co-authors) (hereinafter, "McCullough, *Multifaceted*").

³⁸ Peter A. McCullough et al., Pathophysiological Basis and Rationale for Early Outpatient Treatment of SARS-CoV-2 (COVID-19) Infection, 134 American Journal of Medicine 16, 16 (Jan. 2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7410805/pdf/main.pdf</u> (last visited Oct. 14, 2021) (including 23 co-authors) (hereinafter, "McCullough, Pathophysiological").

³⁹ Physicians Declaration, Global COVID Summit, International Alliance of Physicians and Medical Scientists (Sept. 2021), <u>https://doctorsandscientistsdeclaration.org/</u> (last visited Oct. 14, 2021).

These groups of physicians have established protocols for early treatment, and ivermectin and hydroxychloroquine are staples of those treatments.⁴⁰ As discussed in greater detail below, while the scientific literature is continuing to grow, some data suggest that ivermectin- or hydroxychloroquine-based early treatments of COVID-19 can be effective in thwarting hospitalization and death.⁴¹

3. Ivermectin

A. History of Ivermectin

Researchers discovered ivermectin in the 1970s, and while its first use was to treat parasites in animals, ivermectin has been used in humans since the 1980s.⁴² In the early years, ivermectin effectively stymied the scourge of two devastating parasitic diseases— onchocerciasis (also known as river blindness) and lymphatic filariasis—"among poverty-stricken populations throughout the tropics."⁴³ These are two of the most "disfiguring diseases" that "have plagued the world's poor . . . for centuries."⁴⁴ Later, the use of ivermectin was expanded to include "the treatment of scabies and lice."⁴⁵

43 Id.

Andy Crump & Satoshi Ömura, Ivermectin, 'wonder drug' from Japan: the human use perspective, 87 Proceedings of the Japan Academy, Series B, Physical and biological sciences 13, 13 (2011), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3043740/pdf/pjab-87-013.pdf</u> (last visited Oct. 14, 2021).

E.g., McCullough, Multifaceted, supra, at 519 Table 1 (listing early treatment kits that include both ivermectin and hydroxychloroquine); McCullough, Pathophysiological, supra, at 18–19 (discussing hydroxychloroquine).

⁴¹ E.g., Flavio A. Cadegiani et al., Early COVID-19 therapy with azithromycin plus nitazoxanide, ivermectin or hydroxychloroquine in outpatient settings significantly improved COVID-19 outcomes compared to known outcomes in untreated patients, New Microbes and New Infections (Sept. 2021), available at https://www.sciencedirect.com/science/article/pii/S2052297521000792 (last visited Oct. 14, 2021) (finding that "the use of nitazoxanide, ivermectin[,] and hydroxychloroquine demonstrated unexpected improvements in COVID-19 outcomes when compared to untreated patients").

⁴² Andy Crump, *Ivermectin: enigmatic multifaceted 'wonder' drug continues to surprise and exceed expectations*, 70 The Journal of Antibiotics 495, 495 (2017), *available at <u>https://www.nature.com/articles/</u> <u>ja201711.pdf</u> (last visited Oct. 14, 2021) (hereinafter, "Crump, <i>Ivermectin*").

⁴⁵ Andrew Bryant et al., *Ivermectin for Prevention and Treatment of COVID-19 Infection: A Systematic Review, Meta-analysis, and Trial Sequential Analysis to Inform Clinical Guidelines,* 28 American Journal of Therapeutics 434, 435 (Jul./Aug. 2021), *available at <u>https://journals.lww.com/americantherapeutics/</u> fulltext/2021/08000/ivermectin for prevention and treatment of.7.aspx (last visited Oct. 14, 2021) (hereinafter, "Bryant, <i>Ivermectin*").

Given its track record as a medicine for humans, ivermectin has long since been "approved as an antiparasitic" by the World Health Organization (WHO) and the FDA.⁴⁶ The WHO has also recognized ivermectin as one of its "Essential Medicines.^{*47} Further recognizing the importance of this drug, in 2015 its discoverers won the Nobel Prize in Medicine for their work in uncovering it and bringing it to market.⁴⁸

In the decade leading up to the COVID-19 pandemic, studies began to show ivermectin's surprising versatility. By 2017, ivermectin had "demonstrate[d] antiviral activity against several RNA viruses by blocking the nuclear trafficking of viral proteins."⁴⁹ One recent systematic review cited more than a handful of studies to "demonstrate that ivermectin has antiviral properties against an increasing number of RNA viruses, including influenza, *Zika*, HIV, [and] *Dengue*."⁵⁰ And another review summarized the "antiviral effects of ivermectin" demonstrated through "studies over the past 50 years."⁵¹

Before the pandemic, scholarly literature had also recognized ivermectin's "antiinflammatory capacity."⁵² Doctors thus have been using ivermectin to treat "rosacea, a chronic inflammatory disease," that manifests itself as a reddening of the face, and the FDA has approved ivermectin for that purpose.⁵³ Ivermectin's ability to "curb inflammation," one reviewer wrote, may also "be useful in treating . . . inflammatory airway diseases."⁵⁴ Summing it up, that same reviewer recognized that "ivermectin is continuing

46 Id.

47 Id.

⁴⁸ The Nobel Prize, Press Release for The Nobel Prize in Physiology or Medicine 2015 (Oct. 5, 2015), <u>https://www.nobelprize.org/prizes/medicine/2015/press-release/</u> (last visited Oct. 14, 2021).

49 Crump, Ivermectin, supra, at 500.

⁵⁰ Pierre Kory et al., Review of the Emerging Evidence Demonstrating the Efficacy of Ivermectin in the Prophylaxis and Treatment of COVID-19, 28 American Journal of Therapeutics 299, 301 (2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8088823/</u> (last visited Oct. 14, 2021).

⁵¹ Fatemeh Heidary & Reza Gharebaghi, *Ivermectin: a systematic review from antiviral effects to* COVID-19 complementary regimen, 73 The Journal of Antibiotics 593, 593 (2020), available at <u>https://www.nature.com/articles/s41429-020-0336-z.pdf</u> (last visited Oct. 14, 2021) ("Several studies reported antiviral effects of ivermectin on RNA viruses Furthermore, there are some studies showing antiviral effects of ivermectin against DNA viruses ").

52 Crump, Ivermectin, supra, at 499.

Leon H. Kircik et al., Over 25 Years of Clinical Experience With Ivermectin: An Overview of Safety for an Increasing Number of Indications, 15 Journal of Drugs in Dermatology 325, 325 (Mar. 2016), available at <u>https://jddonline.com/articles/dermatology/S1545961616P0325X</u> (last visited Oct. 14, 2021).

⁵⁴ Crump, Ivermectin, supra, at 499; see also Arianna Portmann-Baracco et al., Antiviral and antiinflammatory properties of ivermectin and its potential use in Covid-19, 56 Archivos De Bronconeumologia

to surprise and excite scientists, offering more and more promise to help improve global public health by treating a diverse range of diseases."55

For more than three decades, ivermectin has also shown itself to be very safe. Indeed, the National Institutes of Health ("NIH") recognize that "ivermectin has been widely used and is generally well tolerated."⁵⁶ One recent systematic review similarly states that "ivermectin at the usual doses . . . is considered extremely safe for use in humans."⁵⁷ Other studies have noted that the medicine "has an established safety profile for human use,"⁵⁸ and it "provide[s] a high margin of safety for a growing number of indications."⁵⁹ Notably, a December 2018 WHO-supported application to add ivermectin as an essential medicine for scabies reviewed the data and concluded that the adverse events associated with ivermectin are "primarily minor and transient."⁶⁰

The available data support this conclusion. The WHO's VigiAccess database, which compiles adverse drug reactions from throughout the world, breaks down the reported side effects for drugs into different categories.⁶¹ The largest reported categories for ivermectin include skin issues, headaches, dizziness, and gastrointestinal disturbances such as diarrhea and nausea.⁶² The NIH confirms that ivermectin's primary adverse side effects "include dizziness, pruritis [itchy skin], nausea, or diarrhea.⁶³ And

⁵⁶ National Institutes of Health, COVID-19 Treatment Guidelines: Ivermectin, <u>https://www.covid19</u> treatmentguidelines.nih.gov/therapies/antiviral-therapy/ivermectin/ (last visited Oct. 14, 2021) (hereinafter, "NIH, COVID-19 and Ivermectin").

57 Bryant, Ivermectin, supra, at 435.

⁵⁸ Leon Caly et al., The FDA-approved drug ivermectin inhibits the replication of SARS-CoV-2 in vitro, Antiviral Research 178 at 3 (June 2020), available at <u>https://www.sciencedirect.com/science/article/pii/S0166354220302011</u> (last visited Oct. 14, 2021).

59 Kircik, Ivermectin, supra, at 325.

⁶⁰ WHO Expert Committee on the Selection and Use of Essential Medicines: Application for inclusion of ivermectin on the WHO Model List of Essential Medicines (EML) and Model List of Essential Medicines for Children (EMLc) for the indication of Scabies at 19 (Dec. 2018), available at https://www.who.int/selection_medicines/committees/expert/22/applications/s6.6 ivermectin.pdf (last visited Oct. 14, 2021).

⁶¹ VigiAccess, Uppsala Monitoring Centre, WHO Collaborating Centre for International Drug Monitoring, <u>http://www.vigiaccess.org/</u> (last visited Oct. 14, 2021).

62 Id.

63 NIH, COVID-19 and Ivermectin, supra-

^{831, 831 (2020),} available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7578741/pdf/main.pdf (last visited Oct. 14, 2021) ("Ivermectin has a demonstrated anti-inflammatory effect in vivo and in vitro").

⁵⁵ Crump, Ivermectin, supra, at 495.

a recent review of ivermectin similarly describes the common side effects as "itching, rash, swollen lymph nodes, joint pain[], fever, and headache."64

The data show not only that the adverse side effects are minor, but also that the percentage of people who report experiencing any adverse events is vanishingly small. The latest statistics available through VigiAccess report only 5,674 adverse drug reactions from ivermectin between 1992 and October 13, 2021.⁶⁵ This number is incredibly low considering that "more than 3.7 billion doses" of ivermectin have been administered to humans worldwide since the 1980s.⁶⁶

To illustrate the safety of ivermectin, compare its VigiAccess report to that of remdesivir, an FDA-approved treatment for COVID-19.⁶⁷ Remdesivir was not released for widespread use until 2020. Yet in the short period of time that it has been on the market, people have reported at least 7,491 adverse drug reactions on VigiAccess, more than ivermectin has registered over the last 30 years.⁶⁸ What's more, serious adverse reactions from remdesivir are reported in high numbers. For example, in less than two years, those who have used remdesivir have reported over 560 deaths, 550 serious cardiac disorders (such as bradycardia and cardiac arrest), and 475 acute kidney injuries.⁶⁹ Since that safety profile is sufficient to retain FDA approval, ivermectin's safety record cannot reasonably be questioned.

B. Ivermectin and COVID-19

As discussed above, ivermectin had shown its antiviral and anti-inflammatory properties long before the pandemic began. So when COVID-19 began to spread across the globe, some in the medical community quickly identified ivermectin as a potential drug for the prevention and treatment of COVID-19. Initially, a group of researchers found that ivermectin significantly inhibited replication of SARS-CoV-2 in cell cultures.⁷⁰ Dismissing

⁶⁵ VigiAccess, Uppsala Monitoring Centre, WHO Collaborating Centre for International Drug Monitoring, <u>http://www.vigiaccess.org/</u> (last visited Oct. 14, 2021).

⁶⁶ Morimasa Yagisawa et al., *Global trends in clinical studies of ivermectin in COVID-19*, 74 The Japanese Journal of Antibiotics 44, 46 (Mar. 2021), *available at <u>http://jja-contents.wdc-jp.com/pdf/JJA74/74-1-open/74-1 44-95.pdf</u> (last visited Oct. 14, 2021).*

⁶⁷ U.S. Food and Drug Administration, FDA Approves First Treatment for COVID-19 (Oct. 22, 2020), https://www.fda.gov/news-events/press-announcements/fda-approves-first-treatment-covid-19 (last visited Oct. 14, 2021).

⁶⁸ VigiAccess, Uppsala Monitoring Centre, WHO Collaborating Centre for International Drug Monitoring, <u>http://www.vigiaccess.org/</u> (last visited Oct. 14, 2021).

69 Id.

70 Caly, supra, at 1.

⁶⁴ Kory, supra, at 314.

that finding, ivermectin doubters argued that too much of the drug would be needed to achieve this antiviral activity in humans.⁷¹ But peer-reviewed models undermined those concerns by showing that the predicted accumulation of ivermectin in the lungs—the site in the body where the medicine is most needed—would be over 10 times higher than necessary for antiviral activity.⁷² In layman's terms, these models indicated that an effective level of the medicine can be reached in lung tissue without creating toxicity in the blood. Plus, other pro-ivermectin doctors have explained that the amount of the drug "required for an effect in cell culture models bear[s] little resemblance to human physiology" because cell cultures lack "an active immune system working synergistically with" the medicine.⁷³

The doctors who believed that ivermectin could be effective against COVID-19 also identified its anti-inflammatory properties as an important countermeasure to the disease. One reason why COVID-19 progresses to its severe phase, many believe, is "the provocation of an overwhelming and injurious inflammatory response."⁷⁴ Thus, ivermectin's anti-inflammatory effects suggest that it can help COVID-19 patients as the disease worsens.

i. Ivermectin Studies and Meta-analyses

Since the COVID-19 pandemic began, researchers have conducted over 20 randomized controlled trials (RCTs) and more observational trials to evaluate ivermectin's effectiveness in the prevention and treatment of COVID-19.⁷⁵ Many of those trials showed promise. On the question of COVID-19 prevention, the Shouman study out of Egypt—a RCT—evaluated ivermectin as a potential prophylaxis for close family members of COVID-19 patients.⁷⁶ The test group included 203 family members who took

74 Id.

⁷¹ Virginia D. Schmith et al., The Approved Dose of Ivermectin Alone is not the Ideal Dose for the Treatment of COVID-19, 108 Clinical Pharmacology & Therapeutics 762, 762 (Oct. 2020), available at https://ascpt.onlinelibrary.wiley.com/doi/epdf/10.1002/cpt.1889 (last visited Oct. 14, 2021).

⁷² Usman Arshad et al., Prioritization of Anti-SARS-Cov-2 Drug Repurposing Opportunities Based on Plasma and Target Site Concentrations Derived from their Established Human Pharmacokinetics, 108 Clinical Pharmacology and Therapeutics 775, 785 (Oct. 2020), available at <u>https://ascpt.onlinelibrary.</u> wiley.com/doi/epdf/10.1002/cpt.1909 (last visited Oct. 14, 2021).

⁷³ Kory, supra, at 301.

⁷⁵ Bryant, Ivermectin, supra, at 435.

⁷⁶ Waheed M. Shouman et al., Use of Ivermectin as a Potential Chemoprophylaxis for COVID-19 in Egypt: A Randomised Clinical Trial, 15 Journal of Clinical and Diagnostic Research 27, 27 (Feb. 2021), available at <u>https://www.jcdr.net/articles/PDF/14529/46795_CE[Ra]_F(Sh)_PF1(SY_OM)_PFA_(OM)_PN(KM).pdf</u> (last visited Oct. 14, 2021).

ivermectin, and only 15 of them (7.4%) developed COVID-19.⁷⁷ Compare that to the 101 family members in the control group, 59 of whom (58.4%) tested positive during the study.⁷⁸ These outcomes prompted the research team to conclude that ivermectin is "a promising, effective[,] and safe chemoprophylactic drug in management of COVID-19.^{*79} Also, the Behera study in India tested ivermectin as a prophylaxis in a group of 3,532 healthcare workers.⁸⁰ Of the 2,199 workers who took two doses of ivermectin prophylaxis three days apart, only 45 (2%) tested positive for COVID-19.⁸¹ But of the 1,147 workers who did not take ivermectin, 133 (11.6%) contracted the disease.⁸² Behera's team thus announced that two doses of ivermectin "as chemoprophylaxis among [healthcare workers] reduced the risk of COVID-19 infection by 83% in the following month.^{*83}

Moving beyond ivermectin's role as a prophylaxis, other studies have demonstrated its potential as a COVID-19 treatment. The Mahmud study—a RCT that explored ivermectin as an early treatment for 363 individuals—concluded that "[p]atients with mildto-moderate COVID-19 infection treated with ivermectin plus doxycycline recovered earlier, were less likely to progress to more serious disease, and were more likely to be COVID-19 negative . . . on day 14.⁸⁴ And Niaee's research team found that ivermectin can help even hospitalized patients.⁸⁵ That group conducted a "randomized, doubleblind, placebo-controlled, multicenter clinical trial" with 180 hospitalized patients diagnosed with COVID-19.⁸⁶ They concluded that ivermectin "reduces the rate of

Priyamadhaba Behera et al., Prophylactic Role of Ivermectin in Severe Acute Respiratory Syndrome Coronavirus 2 Infection Among Healthcare Workers, Cureus, at 1 (Aug. 2021), available at <u>https://assets.cureus.com/uploads/original_article/pdf/64807/20210904-4912-omcmtf.pdf</u> (last visited Oct. 14, 2021).

- a1 Id. at 5.
- 82 Id.
- 83 Id. at 1.

⁸⁴ Reaz Mahmud et al., Ivermectin in combination with doxycycline for treating COVID-19 symptoms: a randomized trial, Journal of International Medical Research 49(5) (Apr. 2021), available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8127799/pdf/10.1177_03000605211013550.pdf (last visited Oct. 14, 2021).

⁶⁵ Morteza Shakhsi Niaee et al., Ivermectin as an adjunct treatment for hospitalized adult COVID-19 patients: A randomized multi-center clinical trial, 14 Asian Pacific Journal of Tropical Medicine 266, 266 (2021), available at <u>https://www.apitm.org/temp/AsianPacJTropMed146266-5371482_145514.pdf</u> (last visited Oct. 14, 2021).

86 Id.

⁷⁷ Id.

⁷⁸ Id.

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⁷⁹ Id.

mortality . . . and duration of hospitalization in adult COVID-19 patients," and "[t]he improvement of other clinical parameters showed that the ivermectin, with a wide margin of safety, had a high therapeutic effect on COVID-19."87

As the data accumulated, scholars began conducting and publishing metaanalyses of the available studies. One such analysis—the Bryant review—focused on 24 total RCTs involving 3,406 participants and found "with moderate certainty that ivermectin treatment in COVID-19 provides a significant survival benefit."⁸⁸ It also concluded that "[u]sing ivermectin early in the clinical course may reduce numbers progressing to severe disease" and that "[t]he apparent safety and low cost suggest that ivermectin is likely to have a significant impact on the SARS-CoV-2 pandemic globally."⁶⁹ Following Bryant's publication of his team's review, the Elgazzar study—one of the RCTs included in the meta-analysis—was questioned and is now under review. This prompted Bryant's team to reanalyze the data without the Elgazzar study, and that review still found "a clear result, showing a 49% reduction in mortality in favor of ivermectin."⁹⁰

Another meta-analysis known as the Popp review has reached more skeptical conclusions. That analysis, which excluded some of the RCTs that Bryant considered, evaluated only 14 studies with 1,678 participants and determined that the "completed studies are small and few are considered high quality."⁹¹ Thus, the authors expressed "uncertain[ty] about the efficacy and safety of ivermectin used to treat or prevent COVID-19."⁹² Recently, however, the Bryant team critiqued the Popp review, highlighting, among other things, that although "Popp claims to provide a 'complete evidence profile,'" it actually "excludes most of the available evidence."⁹³

In further contrast, a third meta-analysis expressed doubt about ivermectin. That one—the Roman review—restricted the pool of RCTs even further, considering only 10

⁹¹ Maria Popp et al., *Ivermectin for preventing and treating COVID-19*, Cochrane Database of Systematic Reviews, at 2 (July 28, 2021), *available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/</u> PMC8406455/pdf/CD015017.pdf (last visited Oct. 14, 2021).*

92 Id.

⁹³ Edmund J. Fordham et al., The uses and abuses of systematic reviews: the case of ivermectin in Covid-19, OSF Preprints, at 7 (Sept. 3, 2021), available at <u>https://osf.io/peqci/</u> (last visited Oct. 14, 2021).

⁸⁷ Id.

Bryant, Ivermectin, supra, at 451.

⁸⁹ Id. at 435.

Andrew Bryant et al., Letter to the Editor: Ivermectin for Prevention and Treatment of COVID-19 Infection: A Systematic Review, Meta-analysis, and Trial Sequential Analysis to Inform Clinical Guidelines, 28 American Journal of Therapeutics 573, 573 (Sept./Oct. 2021), available at <u>https://covid19critical care.com/wp-content/uploads/2021/09/Response-to-Elgazzar.pdf</u> (last visited Oct. 14, 2021).

of them.⁹⁴ After doing this, the authors concluded that ivermectin does "not reduce allcause mortality, [length of hospital stay], or viral clearance . . . in patients with mostly mild COVID-19.⁹⁵ As a result, the researchers announced that ivermectin "is not a viable option to treat patients with COVID-19.⁹⁶

In the days since its publication, the Roman review has drawn some harsh criticism. In particular, the authors of the Bryant review have highlighted four categories of flaws with Roman's work: (1) "mis-reporting of source data," (2) "highly selective study inclusion," (3) "cherry picking' of data within included studies," and (4) "conclusions that do not follow from the evidence."⁹⁷ To illustrate these flaws, consider that Roman's paper initially inverted the treatment and control arms for the Niaee study and thus indicated less mortality in the control group when in fact the opposite was true.⁹⁸ Once that error was fixed, the numbers no longer supported the conclusion that ivermectin does "not reduce all-cause mortality."⁹⁹ Yet the Roman team did not adjust that statement, and thus its "conclusions are no longer based on the data."¹⁰⁰

Furthermore, in a letter to the editor of the American Journal of Therapeutics, two researchers recently explained that Roman's conclusion of no mortality reduction "is not based on the results of the statistical analysis of the data . . . ; instead, it was based on a somewhat vague and possibly biased subjective assessment of the quality of the trials

100 Id.

⁹⁴ Yuani M. Roman et al., Ivermectin for the treatment of Coronavirus Disease 2019: A systematic review and meta-analysis of randomized controlled trials, Clinical Infectious Diseases, at 1 (June 28, 2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8394824/pdf/ciab591.pdf</u> (last visited Oct. 14, 2021).

⁹⁵ Id.

⁹⁶ Id.

⁹⁷ Letter from Andrew Bryant et al. to Robert T. Schooley, MD, Editor in Chief, Clinical Infectious Diseases, at 3, available at <u>https://covid19criticalcare.com/wp-content/uploads/2021/07/RomanRebuttal</u> <u>v7 EF letterhead ML-1.pdf</u> (last visited Oct. 14, 2021) (hereinafter, "Bryant Letter to Schooley").

⁹⁸ Compare Yuani M. Roman et al., *Ivermectin for the treatment of COVID-19: A systematic review* and meta-analysis of randomized controlled trials, Preprint Version 1, at 27 Figure 2 (May 25, 2021), available at <u>https://www.medrxiv.org/content/10.1101/2021.05.21.21257595v1.full.pdf</u> (last visited Oct. 14, 2021) (listing the Niaee study as having four deaths in the control arm and 11 in the ivermectin arm), with Yuani M. Roman et al., *Ivermectin for the treatment of COVID-19: A systematic review and meta-analysis* of randomized controlled trials, Preprint Version 2, at 27 Figure 2 (May 26, 2021), available at <u>https://www.medrxiv.org/content/10.1101/2021.05.21.21257595v2.full.pdf</u> (last visited Oct. 14, 2021) (correcting the Niaee study to list 11 deaths in the control arm and four in the ivermectin arm).

⁹⁹ Bryant Letter to Schooley, supra, at 2.

themselves."¹⁰¹ Those researchers conducted their own Bayesian analysis, a method of statistical inference, and found that the "probability for the hypothesis of a causal link between COVID-19 severity, ivermectin, and mortality is over 99%."¹⁰² As they concluded, "[i]n our view, this Bayesian analysis, based on the statistical study data, provides sufficient confidence that ivermectin is an effective treatment for COVID-19 and this belief supports the conclusions of Bryant over those of Roman."¹⁰³ Those scholars have since published their full analysis in a paper available online.¹⁰⁴

Additional supportive evidence for Bryant's conclusions is a non-peer-reviewed website that currently maintains a running list of 64 COVID-19-related ivermectin studies—RCTs and others—which include all the relevant ivermectin studies except the few (such as Elgazzar) whose data have been called into question.¹⁰⁵ Of those 64 studies, 31 are RCTs and 44 have been peer-reviewed.¹⁰⁶ That site posts multiple metaanalyses of different groupings of the data and concludes that "[m]eta analysis using the most serious outcome reported shows" that ivermectin leads to 66% "improvement for early treatment" and an 86% "improvement for . . . prophylaxis."¹⁰⁷ These "[r]esults are very robust," the site reports, because "in worst case exclusion sensitivity analysis 53 of 64 studies must be excluded to avoid finding statistically significant efficacy."¹⁰⁸

Finally, a recent mini-review of ivermectin and COVID-19 considered the studies analyzing ivermectin's safety specifically in the context of COVID-19 treatments.¹⁰⁹ That mini-review—which was authored by Yale Professor Alessandro D. Santin—observed

102 Id.

¹⁰⁵ Ivermectin for COVID-19: Real-time meta analysis of 64 studies (Oct. 8, 2021), <u>https://ivmmeta.com/</u> (last visited Oct. 14, 2021).

106 Id.

107 Id.

108 Id.

¹⁰⁹ Alessandro D. Santin et al., *Ivermectin: a multifaceted drug of Nobel prize-honoured distinction with indicated efficacy against a new global scourge, COVID-19*, New Microbes New Infections (Aug. 2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8383101/pdf/main.pdf</u> (last visited Oct. 14, 2021).

¹⁰¹ Martin Neil & Norman Fenton, *Bayesian Hypothesis Testing and Hierarchical Modeling of Ivermectin Effectiveness*, 28 American Journal of Therapeutics 576, 576 (Sept/Oct. 2021), *available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8415515/pdf/ajt-28-e576.pdf* (last visited Oct. 14, 2021).

¹⁰³ Id. at 578.

¹⁰⁴ Martin Neil & Norman Fenton, Bayesian hypothesis testing and hierarchical modelling of ivermectin effectiveness in treating Covid-19 (Oct. 1, 2021), available at <u>https://arxiv.org/ftp/arxiv/papers/2109/2109.</u> <u>13739.pdf</u> (last visited Oct. 14, 2021).

that ivermectin "has been safely used in 3.7 billion doses since 1987" and that the medicine has been "used without serious [adverse effects]" in multiple "COVID-19 treatment studies."¹¹⁰

The existing ivermectin studies and meta-analyses are subject to vigorous ongoing disputes, and there are large ongoing studies, at least one of which includes the NIH as a collaborator, that will hopefully provide additional clarity.¹¹¹ But based on the existing medical literature, we do not find clear and convincing evidence that a physician who prescribes ivermectin for COVID-19 after obtaining informed consent engages in unprofessional conduct or otherwise violates the UCA.

While we find the studies and meta-analyses sufficient to resolve this question, we note that epidemiological evidence—derived by analyzing COVID-related data from various states, countries, or regions—is also instructive in the context of a global pandemic. We highlight just a few examples.

One set of scholars analyzed data comparing the COVID-19 rates of countries that routinely administer ivermectin as a prophylaxis and countries that do not.¹¹² The research revealed that "countries with routine mass drug administration of prophylactic . . . ivermectin have a significantly lower incidence of COVID-19."¹¹³ This "highly significant" correlation manifests itself not only "in a worldwide context" but also when comparing African countries that regularly administer prophylactic "ivermectin against parasitic infections" and African countries that do not.¹¹⁴ Based on these results, the researchers surmised that these results "may be connected to ivermectin's ability to inhibit SARS-CoV-2 replication, which likely leads to lower infection rates."¹¹⁵

¹¹² Martin D. Hellwig & Anabela Maia, A COVID-19 prophylaxis? Lower incidence associated with prophylactic administration of ivermectin, International Journal of Antimicrobial Agents (2021), available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7698683/pdf/main.pdf (last visited Oct. 14, 2021).

113 Id. at 1.

114 Id.

115 Id.

¹¹⁰ Id. at 4.

E.g., U.S. National Library of Medicine, ACTIV-6: COVID-19 Study of Repurposed Medications, https://clinicaltrials.gov/ct2/show/NCT04885530?term=activ-6&draw=2&rank=1 (last visited Oct. 14, 2021) (purpose of this trial involving an estimated 15,000 participants is "to evaluate the effectiveness of repurposed medications" that include ivermectin "in reducing symptoms of non-hospitalized participants with mild to moderate COVID-19"); U.S. National Library of Medicine, COVID-OUT: Early Outpatient Treatment for SARS-CoV-2 Infection (COVID-19), <u>https://clinicaltrials.gov/ct2/show/NCT04510194? term=ivermectin+boulware&draw=2&rank=1</u> (last visited Oct. 14, 2021) (purpose of this trial involving 1,160 participants is to understand whether ivermectin is superior to other options, including placebo, in "non-hospitalized adults with SARS-CoV-2 disease for preventing Covid-19 disease progression").

More specifically, Peru's COVID-19 statistics, which have been analyzed in preprint studies and discussed in published ivermectin reviews, are also informative.¹¹⁸ Peru deployed mass ivermectin-based COVID-19 treatments from April 2020 through November 2020 throughout its 25 states.¹¹⁷ In ten of those states, a maximal amount of "mass [ivermectin] treatments of COVID-19 were conducted through a broadside, armyled effort, *Mega-Operación Tayta (MOT)*.^{*118} Fourteen other states had a medium distribution of ivermectin administered at the local level.¹¹⁹ And one state, Lima, distributed a minimal amount of ivermectin due to restrictive government policies.¹²⁰ "The mean reduction in excess deaths 30 days after peak deaths was 74% for the maximal [ivermectin] distribution group, 53% for the medium group[.] and 25% for Lima.^{*121} Furthermore, throughout the country of Peru, "excess deaths decreased 14-fold over four months" leading up to December 1, 2020, "after which deaths then increased 13-fold when [ivermectin] use was restricted under a new president.^{*122}

- 119 Chamie-Quintero, supra, at 2.
- 120 Id.
- 121 Id.
- 122 Id.

¹¹⁶ Juan J. Chamie-Quintero et al., *Ivermectin for COVID-19 in Peru:* 14-fold reduction in nationwide excess deaths, p < 0.002 for effect by state, then 13-fold increase after ivermectin use restricted (Mar. 2021), available at <u>https://osf.io/9egh4/</u> (last visited Oct. 14, 2021); see also Santin, supra, at 3–4 (discussing the Peruvian data); Kory, supra, at 311–13 (same).

¹¹⁷ Chamie-Quintero, supra, at 2.

¹¹⁸ Santin, supra, at 3.

Ivermectin for COVID-19 in Peru: 14-fold reduction in nationwide excess deaths, p=.002 for effect by state, then 13-fold increase after ivermectin use restricted

Juan J. Chamie-Quintero,* Jennifer A. Hibberd,* David E Scheime



Figure 1. A) Excess all-cause deaths (all ages), national population of Peru. These decreased 14-fold August 1 through December 1. 2020; then after IVM use was restricted, increased 13-fold through February 1. All y values are 7-day moving averages: for B.C. ages 2 60. Data are from Peru's National Death Information System (SINADEF).¹² B) Drops in excess deaths for all states of operation *MOT*, an army-led program of mass IVM distributions, but Pasco, which had them on 3 dates. • *MOT* start date: **a** peak deaths: **B** day of peak deaths = 30 days. Jumin also distributed IVM 13 days before *MOT* start. (2) Reductions in excess deaths at -30 days after peak deaths for the 25 states by extent of IVM distributions: maximal-*MOT* (+), mean -74%; moderate-local distributions (c), mean -53%; and minimal-Lima (x). -25%. These reductions for the 25 states correlated with extent of IVM distributions with Kendall to p=0.002.

"Potential confounding factors, including lockdowns and herd immunity, were ruled out using Google community mobility data, seropositivity rates, population densities and geographic distributions of SARS-CoV-2 genetic variations."¹²³ While these figures do not prove causation, they demonstrate a strong correlation between ivermectin use and mortality reductions.

Moving from Peru to India, the government in the State of Uttar Pradesh—a jurisdiction with a population of more than 200 million—"introduced a large-scale 'prophylactic and therapeutic' use of [i]vermectin" that enabled it "to maintain a lower fatality and

123 Santin, supra, at 4.

positivity rate as compared to other states" in India.¹²⁴ As one state official explained, "Uttar Pradesh was the first state in [India] to introduce large-scale prophylactic and therapeutic use of Ivermectin."¹²⁵ The state's health department introduced ivermectin "as prophylaxis for close contacts of [COVID-19] patients" and "health workers," "as well as for the treatment of the patients themselves."¹²⁶ "Despite being [India's] state with the largest population base and a high population density," that state official added, Uttar Pradesh has "maintained a relatively low positivity rate and cases per million of population."¹²⁷ Although these statements from the Uttar Pradesh government do not prove ivermectin's effectiveness, they are informative and worthy of some consideration.

ii. U.S. Public Health Agencies on Ivermectin

Many public health agencies in the United States have now addressed the topic of ivermectin and COVID-19. The NIH has adopted a neutral position, saying that "[t]here is insufficient evidence . . . to recommend either for or against the use of ivermectin for the treatment of COVID-19."¹²⁸ This position, which the NIH adopted in January 2021, overrode its prior stance of "recommend[ing] against the use of ivermectin for the treatment" of COVID-19.¹²⁹ The reason for the change, the NIH recognized, was that "several randomized trials and retrospective cohort studies of ivermectin use in patients with COVID-19 have been published in peer-reviewed journals."¹³⁰ And some of those studies reported positive outcomes, including "shorter time to resolution of disease manifestations that were attributed to COVID-19, greater reduction in inflammatory marker levels, shorter time to viral clearance, [and] lower mortality rates in patients who received ivermectin than in patients who received comparator drugs or placebo."¹³¹ The NIH nevertheless decided not to recommend the use of ivermectin for COVID-19 because other studies suggest "no benefits" and the NIH thought that the available studies

- 125 Id.
- 126 Id.
- 127 Id.
- 128 NIH, COVID-19 and Ivermectin, supra.
- 129 Yagisawa, supra, at 65.
- ¹³⁰ NIH, COVID-19 and Ivermectin, supra.
- 131 Id.

¹²⁴ Maulshree Seth, *Uttar Pradesh government says early use of Ivermectin helped to keep positivity, deaths low,* The Indian Express (May 12, 2021), *available at <u>https://indianexpress.com/article/cities/</u> <u>lucknow/uttar-pradesh-government-says-ivermectin-helped-to-keep-deaths-low-7311786/</u> (last visited Oct. 14, 2021), and <u>https://www.msn.com/en-in/news/other/uttar-pradesh-government-says-early-use-of-ivermectin-helped-to-keep-positivity-deaths-low/ar-BB1gDp5U</u> (last visited Oct. 14, 2021).*

generally suffered from "methodological limitations."¹³² By making a neutral recommendation, the NIH—which is continuing to collaborate on at least one study investigating ivermectin as a treatment for "mild to moderate COVID-19"¹³³—clearly signaled that physicians should use their discretion in deciding whether to treat COVID-19 patients with ivermectin.

Ignoring the NIH's official position, officials within its agencies have sent contradictory messages. On August 29, 2021, Dr. Anthony Fauci, Director of the National Institute of Allergy and Infectious Diseases (NIAID) within the NIH, went on CNN and announced that "there is no clinical evidence" that ivermectin works for the prevention or treatment of COVID-19.¹³⁴ Expanding on that point, he reiterated that "there is no evidence whatsoever" that it works.¹³⁵ Yet this definitive claim directly contradicts the NIH's recognition that "several randomized trials . . . published in peer-reviewed journals" have reported data indicating that ivermectin is effective as a COVID-19 treatment.¹³⁶

The FDA has similarly charted a course of confusion. In March 2021, the FDA posted a webpage entitled "Why You Should Not Use Ivermectin to Treat or Prevent COVID-19."¹³⁷ Although the FDA's concern was stories of some people using the animal form of ivermectin or excessive doses of the human form, the title broadly condemned any use of ivermectin in connection with COVID-19. Yet there was no basis for its sweeping condemnation. Indeed, the FDA itself acknowledged on that very webpage (and continued to do so until the page changed on September 3, 2021) that the agency had *not* even "reviewed data to support use of ivermectin in COVID-19 patients to treat or to prevent COVID-19."¹³⁸ But without reviewing the available data, which had long

132 Id.

¹³³ U.S. National Library of Medicine, ACTIV-6: COVID-19 Study of Repurposed Medications, <u>https://clinicaltrials.gov/ct2/show/NCT04885530?term=activ-6&draw=2&rank=1</u> (last visited Oct. 14, 2021).

¹³⁴ CNN Health, 'Don't do it': Dr. Fauci warns against taking Ivermectin to fight Covid-19 (Aug. 29, 2021), <u>https://edition.cnn.com/videos/health/2021/08/29/dr-anthony-fauci-ivermectin-covid-19-sotu-vpx.cnn</u> (last visited Oct. 14, 2021).

135 Id.

136 NIH, COVID-19 and Ivermectin, supra.

¹³⁷ U.S. Food and Drug Administration, Why You Should Not Use Ivermectin to Treat or Prevent COVID-19 (archived Mar. 5, 2021), <u>https://web.archive.org/web/20210305163946/https://www.fda.gov/</u> <u>consumers/consumer-updates/why-you-should-not-use-ivermectin-treat-or-prevent-covid-19</u> (last visited Oct. 14, 2021) (hereinafter, "FDA, Why You Should Not Use Ivermectin (Mar. 5, 2021)").

¹³⁸ Id.; see also U.S. Food and Drug Administration, Why You Should Not Use Ivermectin to Treat or Prevent COVID-19 (archived Sept. 2, 2021), <u>https://web.archive.org/web/20210902231921/https://www.</u> <u>fda.gov/consumers/consumer-updates/why-you-should-not-use-ivermectin-treat-or-prevent-covid-19</u> (last visited Oct. 14, 2021) (hereinafter, "FDA, Why You Should Not Use Ivermectin (Sept. 2, 2021)").

since been available and accumulating, it is unclear what basis the FDA had for denouncing ivermectin as a treatment or prophylaxis for COVID-19.

On that same webpage, the FDA also declared that "[i]vermectin is not an anti-viral (a drug for treating viruses).^{*139} It did so while another one of its webpages¹⁴⁰ simultaneously cited a study in *Antiviral Research* that identified ivermectin as a medicine "previously shown to have *broad-spectrum anti-viral activity*.^{*141} It is telling that the FDA deleted the line about ivermectin not being "anti-viral" when it amended the first webpage on September 3, 2021.¹⁴²

The FDA has additionally assailed ivermectin's safety by suggesting, though not outright stating, that even a proper dose of human ivermectin might be dangerous when used to treat COVID-19. For example, the FDA announced that "[t]aking a drug for an unapproved use can be very dangerous" and "[t]his is true of ivermectin."¹⁴³ Yet this ignores the fact that, as discussed above, doctors routinely prescribe medicines for offlabel use and that ivermectin is a particularly well-tolerated medicine with an established safety record. Moreover, it is inconsistent for the FDA to imply that ivermectin is dangerous when used to treat COVID-19 while the agency continues to approve remdesivir¹⁴⁴ despite its spottier safety record, as discussed above.

The FDA has also called into question ivermectin's potential effectiveness. When updating the "Why You Should Not Use Ivermectin" webpage on September 3, 2021, the FDA added this entry: "Currently available data do not show ivermectin is effective against COVID-19."¹⁴⁵ But this claim fails to recognize that several RCTs and at least one metaanalysis suggest that ivermectin is effective against COVID-19.

¹⁴² U.S. Food and Drug Administration, Why You Should Not Use Ivermectin to Treat or Prevent COVID-19 (updated Sept. 3, 2021), <u>https://www.fda.gov/consumers/consumer-updates/why-you-should-not-use-ivermectin-treat-or-prevent-covid-19</u> (last visited Oct. 14, 2021) (hereinafter, "FDA, Why You Should Not Use Ivermectin (Sept. 3, 2021)").

143 FDA, Why You Should Not Use Ivermectin (Mar. 5, 2021), supra.

¹⁴⁴ U.S. Food and Drug Administration, FDA Approves First Treatment for COVID-19 (Oct. 22, 2020), <u>https://www.fda.gov/news-events/press-announcements/fda-approves-first-treatment-covid-19</u> (last visited Oct. 14, 2021).

FDA, Why You Should Not Use Ivermectin (Sept. 3 2021), supra.

¹³⁹ FDA, Why You Should Not Use Ivermectin (Mar. 5, 2021), supra.

¹⁴⁰ U.S. Food and Drug Administration, FAQ: COVID-19 and Ivermectin Intended for Animals (Sept. 3, 2021), <u>https://www.fda.gov/animal-veterinary/product-safety-information/faq-covid-19-and-ivermectin-intended-animals</u> (last visited Oct. 14, 2021).

¹⁴¹ Caly, supra, at 1 (emphasis added).

Moreover, a review of the studies on remdesivir makes it difficult to understand why the FDA would condemn the data supporting ivermectin. The NIH reports only five studies testing remdesivir's efficacy against COVID-19.¹⁴⁶ Three of those five studies show *no benefit* from remdesivir, with the largest of those concluding that remdesivir "did not decrease in-hospital mortality in hospitalized patients.^{*147} Even the two remaining studies are far from compelling. One found that "[h]ospitalized patients... who received 5 days of [remdesivir] had better outcomes," but the difference "was of uncertain clinical importance.^{*148} And while the other study indicated that remdesivir "reduced time to clinical recovery" for "patients with severe COVID-19," it also found "[n]o observed benefit ... in patients with mild or moderate COVID-19" and "[n]o statistically significant difference in mortality.^{*149} Beyond that, in September 2021, the Lancet published the results of a large RCT (the DisCoVeRy trial) that found "[n]o clinical benefit ... from the use of remdesivir in patients who were admitted to hospital for COVID-19, were symptomatic for more than 7 days, and required oxygen support.^{*150} The data on ivermectin thus appears at least as strong as the data on remdesivir.

The FDA's most controversial statement on ivermectin came on August 21, 2021, when it posted a link on Twitter to its "Why You Should Not Use Ivermectin" webpage with this message: "You are not a horse. You are not a cow. Seriously, y'all. Stop it."¹⁵¹

- 147 Id.
- 148 Id.
- 149 Id.

¹⁴⁶ National Institutes of Health, Remdesivir: Selected Clinical Data, <u>https://www.covid19treatment</u> <u>guidelines.nih.gov/tables/table-2a/</u> (last visited Oct. 14, 2021).

¹⁵⁰ Florence Ader et al., Remdesivir plus standard of care versus standard of care alone for the treatment of patients admitted to hospital with COVID-19 (DisCoVeRy): a phase 3, randomised, controlled, open-label trial, The Lancet, at 1 (Sept. 14, 2021), available at https://www.thelancet.com/action/ showPdf?pii=S1473-3099%2821%2900485-0 (last visited Oct. 14, 2021).

¹⁵¹ U.S. FDA, Twitter, <u>https://twitter.com/us_fda/status/1429050070243192839</u> (last visited Oct. 14, 2021).



You are not a horse. You are not a cow. Seriously, y'all. Stop it.



Using the Drug ivermectin to treat COVID-19 can be dangerous and even lethal. The FDA has not approved the drug for that purpose. .9 Ida nov

6:57 AM - Aug 21, 2021 - Twitter Web App

S1.9K Retweets 20.6K Quote Tweets 117.7K Likes

This message is troubling not only because it makes light of a serious matter but also because it inaccurately implies that ivermectin is only for horses or cows.

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Despite its attempts to impugn ivermectin, the FDA appears to recognize that doctors may prescribe it for COVID-19. On September 3, 2021, a change in its website makes this clear. The "Why You Should Not Use Ivermectin" webpage originally said that "[i]f you have a prescription for ivermectin for an FDA-approved use, get it from a legitimate source and take it exactly as prescribed."¹⁵² That same sentence now omits the limitation on prescriptions to FDA-approved uses. It says that "[i]f your health care provider writes you an ivermectin prescription, fill it through a legitimate source such as a pharmacy, and take it exactly as prescribed."¹⁵³ This change implicitly acknowledges that ivermectin may be prescribed off-label for COVID-19.

The CDC has followed in the FDA's footsteps of implying that ivermectin is unsafe. On August 26, 2021, the CDC issued an official advisory entitled "Rapid Increase in Ivermectin Prescriptions and Reports of Severe Illness Associated with Use of Products Containing Ivermectin to Prevent or Treat COVID-19.^{*154} Like the FDA, the CDC's

¹⁵² FDA, Why You Should Not Use Ivermectin (Mar. 5, 2021), supra.

¹⁵³ FDA, Why You Should Not Use Ivermectin (Sept. 3, 2021), supra.

¹⁵⁴ Centers for Disease Control and Prevention, Rapid Increase in Ivermectin Prescriptions and Reports of Severe Illness Associated with Use of Products Containing Ivermectin to Prevent or Treat
sweeping title implies that severe illnesses are arising from the prescribed use of human ivermectin to combat COVID-19, but it supplies no data to indicate that human ivermectin in appropriate doses is harming anyone. On the contrary, the CDC's advisory acknow-ledges that the actual concerns arise from the "use of veterinary products not meant for human consumption" and that the reported "[a]dverse effects [are] associated with ivermectin misuse and overdose."¹⁵⁵ The CDC's instructions to the public confirm that its concerns arise from the improper use of ivermectin creams or animal formulas: "Do not swallow ivermectin products that should be used on skin (e.g., lotions and creams) or are not meant for human use, such as veterinary ivermectin products."¹⁵⁶

None of this undermines the use of human ivermectin in proper doses for the treatment or prevention of COVID-19. If anything, the reported uptick in people resorting to animal ivermectin simply reinforces that COVID-19 patients should be encouraged to discuss human ivermectin with their healthcare providers and that those providers should be allowed to consider the available data with their patients. That would be more beneficial for public health than attempting to obscure the demonstrated safety profile of ivermectin.

The media has added to the confusion and misinformation. On August 30, 2021, the New York Times published an article about ivermectin stating that "Mississippi's health department said earlier this month that *70 percent* of recent calls to the state poison control center had come from people who ingested ivermectin from livestock supply stores."¹⁵⁷ Yet two weeks later, on September 13, 2021, the Times amended its story by deleting that sentence and adding this note after the article: "An earlier version of this article misstated the percentage of recent calls to the Mississippi poison control center related to ivermectin. It was 2 percent, not 70 percent."¹⁵⁸

Similarly, on September 3, 2021, Rolling Stone published a story entitled "Gunshot Victims Left Waiting as Horse Dewormer Overdoses Overwhelm Oklahoma Hospitals,

COVID-19, Health Advisory, at 1 (Aug. 26, 2021), available at https://emergency.cdc.gov/han/2021/pdf/CDC HAN 449.pdf (last visited Oct. 14, 2021).

¹⁵⁵ Id.

¹⁵⁶ Id. at 3.

¹⁵⁷ Emma Goldberg, Demand Surges for Deworming Drug for Covid, Despite No Evidence It Works, New York Times (Aug. 30, 2021), available at <u>https://web.archive.org/web/20210830091038/</u> <u>https://www.nytimes.com/2021/08/30/health/covid-ivermectin-prescriptions.html</u> (last visited Oct. 14, 2021) (emphasis added).

¹⁵⁸ Emma Goldberg, Demand Surges for Deworming Drug for Covid, Despite No Evidence It Works, New York Times (amended Sept. 28, 2021), available at <u>https://www.nytimes.com/2021/08/30/health/covid-ivermectin-prescriptions.html</u> (last visited Oct. 14, 2021).

Doctor Says."¹⁵⁹ Soon thereafter, one the hospitals where this doctor supposedly works denied that claim, and "the doctor [did] not respond[] to requests for further comment."¹⁶⁰ Rather than delete the article or substantially rewrite it, Rolling Stone left the article largely unchanged and amended the title to say: "One Hospital Denies Oklahoma Doctor's Story of Ivermectin Overdoses Causing ER Delays for Gunshot Victims."¹⁶¹ In addition, the magazine added an "update" message stating, among other things, that "[o]ne hospital has denied [the doctor's] claim that ivermectin overdoses are causing emergency room backlogs and delays in medical care in rural Oklahoma, and Rolling Stone has been unable to independently verify any such cases as of the time of this update."¹⁶² In other words, the publication allowed a story based on a discredited and nonresponsive source to remain available to the public. It is no wonder that some people are unsure what to believe about ivermectin.

iii. Foreign Public Health Agencies on Ivermectin

Looking abroad, in March 2021, the WHO "recommend[ed] not to use ivermectin in patients with COVID-19 except in the context of a clinical trial."¹⁶³ The basis for this recommendation rested not on proof that ivermectin is ineffective, but on the WHO's belief that the existing studies were of too low quality to support any conclusive determinations.¹⁶⁴ Notably, though, while the WHO questioned the quality of the evidence, its analysis determined, based on data from 1,419 patients in seven studies, that patients treated with ivermectin had a 14 per 1,000 chance of death while patients in the control groups had a 70 per 1,000 chance of death.¹⁶⁵ Also, the WHO considered only

161 Id.

162 Id.

164 Id.

165 Id. at 23.

Peter Wade, Gunshot Victims Left Waiting as Horse Dewormer Overdoses Overwhelm Oklahoma Hospitals, Doctor Says, Rolling Stone (Sept. 3, 2021), available at <u>https://web.archive.org/web/</u> 20210903231939/https://www.rollingstone.com/politics/politics-news/gunshot-victims-horse-dewormerivermectin-oklahoma-hospitals-covid-1220608/ (last visited Oct. 14, 2021).

Peter Wade, One Hospital Denies Oklahoma Doctor's Story of Ivermectin Overdoses Causing ER Delays for Gunshot Victims, Rolling Stone (amended Sept. 5, 2021), available at <u>https://www.rollingstone.</u> <u>com/politics/politics-news/gunshot-victims-horse-dewormer-ivermectin-oklahoma-hospitals-covid-1220608/</u> (last visited Oct. 14, 2021).

¹⁶³ World Health Organization, Therapeutics and COVID-19: Living Guideline, at 20 (July 6, 2021), available at <u>https://files.magicapp.org/guideline/a6e3f83e-bff5-481c-90ab-130aa86bbe83/published</u> guideline 5486-6 1.pdf (last visited Oct. 14, 2021) (hereinafter, "WHO COVID-19 Guidelines").

ivermectin's effectiveness as a COVID-19 treatment and did not assess its potential as a prophylaxis.¹⁶⁶

Public health authorities in other countries have declined to follow the WHO's guidance. Most importantly, the NIH continues to embrace its neutral recommendation on ivermectin. Also, in May 2021, the State of Goa in India announced, through its health minister Vishwajit Rane, that "it would give [ivermectin] to all its adult residents" in its efforts to combat COVID-19.¹⁶⁷ Likewise, as discussed above, India's Uttar Pradesh continues to distribute ivermectin to people diagnosed with COVID-19. And El Salvador's Ministry of Public Health has included ivermectin as part of its recommendations for early COVID-19 treatment via home patient kit.¹⁶⁸ We did not conduct an exhaustive search on other countries' practices, so this list is simply intended to be illustrative.

iv. Professional Associations and Physicians on Ivermectin

Professional associations, both here in the United States and abroad, have adopted conflicting positions on ivermectin and COVID-19. The American Medical Association (AMA), American Pharmacists Association (APhA), and American Society of Health-System Pharmacists (ASHP) have issued a statement that "strongly oppose[s] the ordering, prescribing, or dispensing of ivermectin to prevent or treat COVID-19 outside of a clinical trial."¹⁶⁹ But this statement relies solely on the FDA's and CDC's statements. Consider the AMA, APhA, and ASHP's claim that "[u]se of ivermectin for the prevention and treatment of COVID-19 has been demonstrated to be harmful to patients."¹⁷⁰ Their only support for that alarming statement is the CDC Health Alert discussed above.¹⁷¹ But as we explained, that CDC advisory gave no indication that any severe adverse effects are occurring from the use of human ivermectin in appropriate doses.

170 Id.

¹⁶⁶ *Id.* at 18.

¹⁶⁷ Siladitya Ray, Indian State Will Offer Ivermectin To Entire Adult Population — Even As WHO Warns Against Its Use As Covid-19 Treatment, Forbes (May 11, 2021), available at <u>https://www.forbes.com/sites/ siladityaray/2021/05/11/indian-state-will-offer-ivermectin-to-entire-adult-population---even-as-who-warnsagainst-its-use-as-covid-19-treatment/?sh=3d45adce6d9f (last visited Oct. 14, 2021).</u>

¹⁶⁸ El Salvador Minister of Public Health Includes Ivermectin as COVID-19 Pandemic Continues, TrialSite News (Aug. 26, 2021), available at <u>https://trialsitenews.com/el-salvador-minister-of-public-health-includes-ivermectin-as-covid-19-pandemic-continues/</u> (last visited Oct. 14, 2021).

¹⁶⁹ American Medical Association, AMA, APhA, ASHP statement on ending use of ivermectin to treat COVID-19 (Sept. 1, 2021), available at <u>https://www.ama-assn.org/press-center/press-releases/ama-aphaashp-statement-ending-use-ivermectin-treat-covid-19</u> (last visited Oct. 14, 2021) (hereinafter, "AMA, APhA, and ASHP Statement on Ivermectin").

Those groups' opposition to ivermectin also conflicts with their otherwise steadfast support for healthcare providers' rights to prescribe medicines for off-label use. They call for ivermectin's ban because the FDA has not approved it "to prevent or treat COVID-19" and some public-health agencies have found "insufficient evidence" to support its use.¹⁷² But just last year, these same professional associations, when discussing prescriptions for hydroxychloroquine to treat COVID-19, affirmed that "[n]ovel off-label use of FDA-approved medications is a matter for the physician's or other prescriber's professional judgment.^{*173} Moreover, the AMA elsewhere recognizes "its strong support for the autonomous clinical decision-making authority of . . . physician[s]" to "lawfully use an FDA approved drug product . . . for an off-label indication when such use is based upon sound scientific evidence.^{*174} In their recent ivermectin statement, however, the AMA, APhA, and ASHP ignore that some sound scientific evidence, including meta-analyses of RCTs, supports the use of ivermectin for COVID-19.

The AMA, APhA, and ASHP mentioned the statement of Merck—the original patentholder on ivermectin—as an additional basis for their position.¹⁷⁵ Yet that does not provide persuasive support for their opposition to ivermectin. Merck's February 2021 statement expressed its view that there is "[n]o meaningful evidence for . . . clinical efficacy in patients with COVID-19,"¹⁷⁶ but this simply ignores the RCTs demonstrating ivermectin's efficacy. Merck then claimed that there is "[a] concerning lack of safety data in the majority of studies."¹⁷⁷ While worded vaguely, this statement, when read carefully, says next to nothing. It simply acknowledges that many of the studies it references did not track safety data. It is not saying, though it might be implying, that the studies showed the medicine to be dangerous. But Merck, of all sources, knows that ivermectin is exceedingly safe, so the absence of safety data in recent studies should not be concerning to the company.

¹⁷² Id.

¹⁷³ American Medical Association, Joint statement on ordering, prescribing or dispensing COVID-19 medications (Apr. 17, 2020), available at <u>https://www.ama-assn.org/delivering-care/public-health/joint-statement-ordering-prescribing-or-dispensing-covid-19</u> (last visited Oct. 14, 2021).

¹⁷⁴ American Medical Association, Patient Access to Treatments Prescribed by Their Physicians, <u>https://policysearch.ama-assn.org/policyfinder/detail/Patient%20Access%20to%20Treatments%20</u> <u>Prescribed%20by%20Their%20Physicians%20H-120.988%20%20?uri=%2FAMADoc%2FHOD.xml-0-201.xml</u> (last visited Oct. 14, 2021).

¹⁷⁵ AMA, APhA, and ASHP Statement on Ivermectin, supra.

¹⁷⁶ Merck, Merck Statement on Ivermectin use During the COVID-19 Pandemic (Feb. 4, 2021), <u>https://www.merck.com/news/merck-statement-on-ivermectin-use-during-the-covid-19-pandemic/</u> (last visited Oct. 14, 2021).

Why would ivermectin's original patentholder go out of its way to question this medicine by creating the impression that it might not be safe? There are at least two plausible reasons. First, ivermectin is no longer under patent, so Merck does not profit from it anymore. That likely explains why Merck declined to "conduct[] clinical trials" on ivermectin and COVID-19 when given the chance.¹⁷⁸ Second, Merck has a significant financial interest in the medical profession rejecting ivermectin as an early treatment for COVID-19. "[T]he U.S. government has agreed to pay [Merck] about \$1.2 billion for 1.7 million courses of its experimental COVID-19 treatment, if it is proven to work in an ongoing large trial and authorized by U.S. regulators."¹⁷⁹ That treatment, known as "molnupiravir, aims to stop COVID-19 from progressing and can be given early in the course of the disease."¹⁸⁰ On October 1, 2021, Merck announced that preliminary studies indicate that molnupiravir "reduced hospitalizations and deaths by half,"¹⁸¹ and that same day its stock price "jumped as much as 12.3%."¹⁸² Thus, if low-cost ivermectin works better than—or even the same as—molnupiravir, that could cost Merck billions of dollars.

While one side of the "professional associations" ledger includes the AMA, APhA, and ASHP (with Merck's backing), other associations disagree with their stance. In particular, the Association of American Physicians and Surgeons (AAPS)—a long-established group that has represented doctors in all specialties since 1943—has raised questions concerning those associations' "startling and unprecedented position that American physicians should immediately stop prescribing, and pharmacists should stop honoring their prescriptions for ivermectin for COVID-19 patients."¹⁸³ The AAPS pointed "out that many physicians disagree with the AMA, writing around 88,000 ivermectin

¹⁷⁸ Yagisawa, supra, at 61.

U.S. signs \$1.2 bln deal for 1.7 mln courses of Merck's experimental COVID-19 drug, Reuters (Jun. 9, 2021), available at <u>https://www.reuters.com/business/healthcare-pharmaceuticals/merck-says-us-govt-buy-about-17-mln-courses-cos-covid-19-drug-2021-06-09/</u> (last visited Oct. 14, 2021).

Matthew Perrone, Merck says COVID-19 pill cuts risk of death, hospitalization, Associated Press (Oct. 1, 2021), available at <u>https://apnews.com/article/merck-says-experimental-covid-pill-cuts-worsteffects-a9a2245fdcee324f6bbd776a0fffcc60</u> (last visited Oct. 14, 2021).

Lewis Krauskopf & Manojna Maddipatia, Merck COVID-19 pill success slams Moderna shares, shakes up healthcare sector, Reuters (Oct. 1, 2021), available at <u>https://www.reuters.com/business/</u><u>healthcare-pharmaceuticals/merck-covid-19-pill-success-slams-moderna-shares-shakes-up-healthcare-sector-2021-10-01/</u> (last visited Oct. 14, 2021).

Association of American Physicians and Surgeons, AAPS Challenges the AMA on Efforts to Suppress Ivermectin Use in COVID (Sept. 4, 2021), available at <u>https://aapsonline.org/aaps-challenges-the-ama-on-efforts-to-suppress-ivermectin-use-in-covid/</u> (last visited Oct. 14, 2021).

prescriptions per week."184 The AAPS has thus publicly resisted these groups' call to "stop[] the off-label use of long-approved drugs."185

In addition, the Tokyo Metropolitan Medical Association, as explained by its chairman Haruo Ozaki, recommended the use of ivermectin for COVID-19 patients in February 2021.¹⁸⁶ That organization emphasized that ivermectin should be administered to people diagnosed with COVID-19 because, among other reasons, it has been effective when used in other countries.¹⁸⁷ Other doctors' groups similarly advocate for ivermectin as a staple of early COVID-19 treatment. The Front Line COVID-19 Critical Care Alliance has been an outspoken supporter. Its organization "regard[s] ivermectin as a core medication in the prevention and treatment of COVID-19,^{*188} and it includes a five-day course of ivermectin as part of its COVID-19 early treatment protocol.¹⁸⁹ Also, the British Ivermectin Recommendation Development Group (BIRD) is a UK-based association of "clinicians, health researchers[,] and patient representatives from all around the world" that collectively "advocate[s] for the use of ivermectin" against COVID-19.¹⁹⁰

In summary, the evidence discussed above shows (1) that ivermectin has demonstrated some effectiveness in preventing and treating COVID-19 and (2) that its side effects are primarily minor and transient. Thus, the UCA does not preclude physicians from considering ivermectin for the prevention or treatment of COVID-19.

184 Id.

185 Id.

187 Id.

188 Front Line COVID-19 Critical Care Alliance, Ivermectin in COVID-19, <u>https://covid19criticalcare.</u> com/ivermectin-in-covid-19/ (last visited Oct. 14, 2021).

¹⁸⁹ Front Line COVID-19 Critical Care Alliance, Prevention & Treatment Protocols for COVID-19, <u>https://covid19criticalcare.com/wp-content/uploads/2020/11/FLCCC-Alliance-I-MASKplus-Protocol-ENGLISH.pdf</u> (last visited Oct. 14, 2021).

¹⁹⁰ British Ivermectin Recommendation Development Group, Who are the BIRD Group, <u>https://bird-group.org/who-are-bird/</u> (last visited Oct. 14, 2021).

¹⁸⁶ Tokyo Metropolitan Medical Association recommends ivermectin administration to prevent aggravation, Nikkei (Feb. 9, 2021), <u>https://www.nikkei.com/article/DGXZQOFB25AAL0V20C21A1000000/</u> (last visited Oct. 14, 2021).

4. Hydroxychloroquine

A. <u>History of Hydroxychloroquine</u>

Hydroxychloroquine, a less toxic derivative of a medicine named chloroquine, was first developed in 1946¹⁹¹ and approved by the FDA in 1955.¹⁹² Since that time, hydroxychloroquine has been widely used as a prophylaxis and treatment for malaria.¹⁹³ It has also "prove[n] to be effective in a number of autoimmune diseases," including systemic lupus erythematosus,¹⁹⁴ primary Sjögren's syndrome, and rheumatoid arthritis, and for those uses, it is often taken daily for years at a time.¹⁹⁵ Hydroxychloroquine's success against these autoimmune diseases "is linked to its anti-inflammatory and immunomodulatory effects."¹⁹⁶ Because of its versatility and efficacy, "[m]illions of hydroxychloroquine doses are prescribed annually."¹⁹⁷ In just the year 2019, hydroxy-chloroquine was prescribed over 5.4 million times in the United States alone.¹⁹⁸

In 2004, long before the COVID-19 pandemic began, a lab study revealed that chloroquine is "an effective inhibitor of the replication of the severe acute respiratory syndrome coronavirus (SARS-CoV) in vitro" and thus that it should "be considered for immediate use in the prevention and treatment of SARS-CoV infections."¹⁹⁹ The following

193 Id.

 ¹⁹⁴ Claudio Ponticelli & Gabriella Moroni, Hydroxychloroquine in systemic lupus erythematosus (SLE),
16 Expert Opinion on Drug Safety 411, 411 (2017), available at <u>https://www.tandfonline.com/</u> doi/full/10.1080/14740338.2017.1269168?scroll=top&needAccess=true (last visited Oct. 14, 2021).

¹⁹⁵ Eliise Laura Nirk et al., Hydroxychloroquine in rheumatic autoimmune disorders and beyond, EMBO Molecular Medicine, at 1 (Aug. 2020), available at <u>https://www.embopress.org/doi/epdf/10.15252/emmm.</u> 202012476 (last visited Oct. 14, 2021).

196 Id.

197 Fram, supra, at 389.

¹⁹⁸ ClinCalc, Hydroxychloroquine Drug Usage Statistics, United States, 2013–2019, <u>https:// clincalc.com/DrugStats/Drugs/Hydroxychloroquine</u> (last visited Oct. 14, 2021).

Els Keyaerts et al., In vitro inhibition of severe acute respiratory syndrome coronavirus by chloroquine, 323 Biochemical and Biophysical Research Communications 264, 264 (2004), available at https://www.sciencedirect.com/science/article/pii/S0006291X0401839X (last visited Oct. 14, 2021).

¹⁹¹ National Institutes of Health, COVID-19 Treatment Guidelines: Chloroquine or Hydroxychloroquine and/or Azithromycin, <u>https://www.covid19treatmentguidelines.nih.gov/therapies/antiviraltherapy/chloroquine-or-hydroxychloroquine-and-or-azithromycin/</u> (last visited Oct. 14, 2021) (hereinafter, "NIH, COVID-19 and Hydroxychloroquine").

¹⁹² Georgi Fram et al., Cardiac Complications Attributed to Hydroxychloroquine: A Systematic Review of the Literature Pre-COVID-19, 17 Current Cardiology Reviews 389, 389 (2021), available at https://www.eurekaselect.com/186876/article (last visited Oct. 14, 2021).

year, another paper explained that "chloroquine has strong antiviral effects on SARS-CoV infection" and "is effective in preventing the spread of SARS[-]CoV in cell culture."200

It is widely recognized in the medical community that hydroxychloroquine is generally safe, so safe in fact that it may be prescribed to pregnant women²⁰¹ and "children of all ages."²⁰² During the beginning of the pandemic, the FDA Commissioner stated that hydroxychloroquine has "a well-established safety profile" for malaria, lupus, and rheumatoid arthritis.²⁰³ According to the CDC, hydroxychloroquine's "most common adverse reactions reported" are minor issues such as "stomach pain, nausea, vomiting, . . . headache," and "itching."²⁰⁴ While the CDC recognizes that high doses, "such as those used to treat rheumatoid arthritis, have been associated with retinopathy," a serious eye condition, that side effect is "extremely unlikely" when hydroxychloroquine is used in short durations with moderate doses.²⁰⁵ Notably, the CDC's guidance on hydroxychloroquine does not mention any concerns about cardiac disorders stemming from the drug.

B. <u>Hydroxychloroquine and COVID-19</u>

At the outset of the pandemic, researchers found—consistent with the prior studies demonstrating chloroquine's efficacy against SARS-CoV—that hydroxychloroquine "can efficiently inhibit SARS-CoV-2 infection in vitro."²⁰⁶ These COVID-19 studies specifically

²⁰² Centers for Disease Control and Prevention, Medicines for the Prevention of Malaria While Traveling Hydroxychloroquine (Plaquenil™), <u>https://www.cdc.gov/malaria/resources/pdf/</u> <u>fsp/drugs/Hydroxychloroquine.pdf</u> (last visited Oct. 14, 2021) (hereinafter, "CDC, Malaria Travel").

U.S. Food & Drug Administration, Bringing a Cancer Doctor's Perspective to FDA's Response to the COVID-19 Pandemic (Mar. 29, 2020), <u>https://www.fda.gov/news-events/fda-voices/bringing-cancerdoctors-perspective-fdas-response-covid-19-pandemic</u> (last visited Oct. 14, 2021) (hereinafter, "FDA, Bringing Perspective").

204 CDC, Malaria Travel, supra.

²⁰⁵ Centers for Disease Control and Prevention, Yellow Book, Chapter 4: Travel-Related Infectious Diseases – Malaria (2020), available at <u>https://wwwnc.cdc.gov/travel/yellowbook/2020/travel-related-infectious-diseases/malaria#1939</u> (last visited Oct. 14, 2021).

Jia Liu et al., Hydroxychloroquine, a less toxic derivative of chloroquine, is effective in inhibiting SARS-CoV-2 infection in vitro, Cell Discovery, at 4 (2020), available at <u>https://www.nature.com/articles/</u> <u>s41421-020-0156-0.pdf</u> (last visited Oct. 14, 2021).

Martin J. Vincent et al., Chloroquine is a potent inhibitor of SARS coronavirus infection and spread, Virology Journal, at 1 (Aug. 2005), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1232869/</u> pdf/1743-422X-2-69.pdf (last visited Oct. 14, 2021).

²⁰¹ Ponticelli & Moroni, supra, at 411; see also Ewa Haładyj et al., Antimalarials - are they effective and safe in rheumatic diseases? 56 Reumatologia 164, 171–72 (2018), available at <u>https://www.ncbi.nlm. nih.gov/pmc/articles/PMC6052376/pdf/RU-56-33240.pdf</u> (last visited Oct. 14, 2021) (noting that hydroxychloroquine "can be continued in the treatment of rheumatic diseases during pregnancy and lactation").

showed that hydroxychloroquine "can inhibit [SARS-CoV-2] virus entry, transmission[,] and replication."²⁰⁷ In addition to this "antiviral activity," hydroxychloroquine also has "anti-inflammatory properties" that help regulate "pro inflammatory cytokines."²⁰⁸ These characteristics—both the antiviral properties and the anti-inflammatory activity—are important countermeasures against COVID-19.

i. Hydroxychloroquine Studies and Meta-analyses

Many large observational studies suggest that hydroxychloroquine significantly reduces the risk of hospitalization and death when administered to outpatients—particularly high-risk outpatients—as part of early COVID-19 treatment. For example, the Mokhtari study "was a multicenter, population-based national retrospective-cohort investigation of 28,759 adults with mild COVID-19 seen . . . between March and September 2020 throughout Iran."²⁰⁹ The data showed that "[t]he odds of hospitalization . . . reduced by 38%" and the chance of death decreased by 73% for those who took hydroxychloroquine.²¹⁰ Critically, those "effects were maintained after adjusting for age, comorbidities, and diagnostic modality," and "[n]o serious [hydroxychloroquine]-related adverse drug reactions were reported."²¹¹

In the same vein, the recently published Million study evaluated 10,429 "adult outpatients" in France infected with SARS-CoV-2 who were "treated early" with hydroxychloroquine plus azithromycin.²¹² Only five deaths occurred among the 8,315 patients who received hydroxychloroquine plus azithromycin—a mere 0.6 per 1,000 patients while 11 died among the 2,114 who received either no treatment or azithromycin alone a much higher rate of 5.2 per 1,000 patients.²¹³ Based on these figures, the study's authors found that hydroxychloroquine "was associated with a lower risk of death, independently of age, sex[,] and epidemic period."²¹⁴ Million's team thus concluded that

208 Id.

210 Id.

211 Id.

²¹² Million, supra, at 1063.

²⁰⁷ Jyoti Bajpai et al., *Hydroxychloroquine and COVID-19 - A narrative review*, 67 Indian Journal of Tuberculosis 147, 148 (Dec. 2020), *available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7836863/</u> pdf/main.pdf (last visited Oct. 14, 2021).*

²⁰⁹ Majid Mokhtari et al., *Clinical outcomes of patients with mild COVID-19 following treatment with hydroxychloroquine in an outpatient setting*, International Immunopharmacology, at 1 (Jul. 2021), *available at* <u>https://www.sciencedirect.com/science/article/pii/S1567576921002721</u> (last visited Oct. 14, 2021).

²¹³ Id. at 1066.

²¹⁴ Id. at 1063.

"[e]arly ambulatory treatment of COVID-19" with hydroxychloroquine plus azithromycin "is associated with very low mortality" and it "improve[s] COVID-19 survival compared to other regimens."²¹⁵

Another group of researchers assessed an elderly population living in a nursing home in the small European state of Andorra.²¹⁶ Their study included "100 COVID-19 confirmed cases" in the nursing home "from March 15 to June 5, 2020."²¹⁷ After evaluating the numbers, these researchers concluded that "[t]reatment with hydroxychloroquine and azithromycin was associated with lower mortality in these patients."²¹⁸ And "the multivariate logistic regression analysis identified hydroxychloroquine plus azithromycin treatment as an independent factor favoring survival compared with no treatment or other treatments."²¹⁹ The study also reinforced hydroxychloroquine's longstanding safety profile because "[c]ardiac monitoring was performed by electrocardiogram, and no rhythm changes were observed . . . in any patient."²²⁰

Added to all this, a preprint of another large observational study by Sulaiman supports the use of hydroxychloroquine as part of early COVID-19 treatment.²²¹ This "study took place in 238 ambulatory fever clinics in Saudi Arabia" during June 2020.²²² Of the 5,541 participating patients, 1,817 were given hydroxychloroquine, and 3,724 received only supportive care.²²³ The researchers found that early hydroxychloroquine-based "therapy was associated with a lower hospital admission" of 9.4% compared to 16.6% for supportive care alone, which equated to a relative risk reduction of 43%. "Adjusting for age, gender, and major comorbid conditions, a multivariate logistic regression model" further confirmed the significant decrease in the hospitalization risk of

- 217 Id.
- 218 Id.
- 219 Id. at 606.
- 220 Id. at 603.

²²¹ Tarek Sulaiman et al., *The Effect of Early Hydroxychloroquine-based Therapy in COVID-19* Patients in Ambulatory Care Settings: A Nationwide Prospective Cohort Study, Preprint, at 1 (2020), available at <u>https://www.medrxiv.org/content/10.1101/2020.09.09.20184143v1.full.pdf</u> (last visited Oct. 14, 2021).

- 222 Id.
- 223 Id.

²¹⁵ Id.

²¹⁶ Eva Heras et al., COVID-19 mortality risk factors in older people in a long-term care center, 12 European Geriatric Medicine 601, 601 (2021), available at <u>https://link.springer.com/content/pdf/10.1007/</u> <u>s41999-020-00432-w.pdf</u> (last visited Oct. 14, 2021).

patients who received hydroxychloroquine.²²⁴ Regression analysis also demonstrated that hydroxychloroquine reduced the mortality risk by an odds ratio of .36, which equates to a threefold drop in deaths.²²⁵ Other observational studies further suggest that hydroxychloroquine has value as an early COVID-19 treatment.²²⁶

We acknowledge that other studies and meta-analyses have concluded that hydroxychloroquine has little to no effect on COVID-19.²²⁷ Yet those materials generally blur the important distinction between hydroxychloroquine's efficacy as an early treatment for mild COVID-19 in nonhospitalized patients and its efficacy as a late treatment for severe COVID-19 in hospitalized patients.²²⁸ As explained above, COVID-19 in its early stages, which consists primarily of cold- and flu-like symptoms, is very different from severe COVID-19, which is a lower respiratory disease often accompanied by respiratory failure and multiple organ dysfunction. Thus, evidence about hydroxychloroquine's use "in inpatients[] is irrelevant with regard to the efficacy of [the drug] in early high-risk outpatient disease."²²⁹ So even if hydroxychloroquine is not effective against severe COVID-19, that does not disprove its value as an early treatment against the disease.

The key, then, is to focus on data that assess hydroxychloroquine's effectiveness in early treatment. A prime example of that is a recently published meta-analysis that combined the Million, Mokhtari, and Sulaiman studies discussed above with two other

²²⁷ Tawanda Chivese et al., Efficacy of chloroquine and hydroxychloroquine in treating COVID-19 infection: A meta-review of systematic reviews and an updated meta-analysis, Travel Medicine and Infectious Disease, at 1 (Sept./Oct. 2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/</u> <u>PMC8273040/pdf/main.pdf</u> (last visited Oct. 14, 2021) (concluding that hydroxychloroquine is "not effective in treating COVID-19").

228 Id. at 3 (noting that this meta-analysis considered studies of people with "confirmed COVID-19, regardless of . . . the severity of illness").

Harvey A. Risch, Early Outpatient Treatment of Symptomatic, High-Risk COVID-19 Patients That Should Be Ramped Up Immediately as Key to the Pandemic Crisis, 189 American Journal of Epidemiology 1218, 1218 (Nov. 2020), available at <u>https://academic.oup.com/aje/article/189/11/1218/5847586</u> (last visited Oct. 14, 2021).

²²⁴ Id.

²²⁵ Id. at 14.

E.g., Andrew Ip et al., Hydroxychloroquine in the treatment of outpatients with mildly symptomatic COVID-19: a multi-center observational study, BMC Infectious Diseases (2021), available at https://bmcinfectdis.biomedcentral.com/track/pdf/10.1186/s12879-021-05773-w.pdf (concluding in a study of 1,274 outpatients with SARS-CoV-2 infection that "there was an association between exposure to hydroxychloroquine and a decreased rate of hospitalization from COVID-19"); Yi Su, Efficacy of early hydroxychloroquine treatment in preventing COVID-19 pneumonia aggravation, the experience from Shanghai, China, 14 BioScience Trends 408, 408 (2020), available at https://bmcinfectdis.biomedcentral.com/track/pdf/10.1186/s12879-021-05773-w.pdf (concluding in a study of 1,274 outpatients with SARS-CoV-2 infection that "there was an association between exposure to hydroxychloroquine treatment in preventing COVID-19 pneumonia aggravation, the experience from Shanghai, China, 14 BioScience Trends 408, 408 (2020), available at https://www.jstage.jst.go.jp/article/bst/14/6/14 2020.03340/ pdf/-char/en (last visited Oct. 14, 2021) (finding in a study of 616 individuals that "[I]he early use of hydroxychloroquine decreased the improvement time and the duration of COVID-19 detection in throat and stool swabs").

outpatient studies.²³⁰ Those five studies together included 32,124 total outpatients, and the analysis revealed that hydroxychloroquine is associated with a 69% reduction in mortality when used as an early COVID-19 treatment.²³¹ In addition, a few months ago, another team of researchers reviewed "nine reports of early treatment outcomes in COVID-19 nursing home patients.^{*232} Data from those studies revealed that "hydroxychloroquine-based multidrug regimens were associated with a statistically significant > 60% reduction in mortality.^{*233} And another scholar, Dr. Harvey A. Risch, Professor of Epidemiology at Yale School of Public Health, has published online a non-peer-reviewed meta-analysis of ten studies exploring hydroxychloroquine as an early COVID-19 treatment.²³⁴ He concluded that for people receiving that treatment the odds ratio of hospitalization was .56 and the odds ratio of death was .25. In other words, his meta-analysis demonstrated that when hydroxychloroquine is administered as an early COVID-19 treatment, it can reduce the risk of death by 75%.

To be sure, these data derive from large-scale observational studies rather than RCTs, and we understand that RCTs are considered the gold standard in medicine. But for at least two reasons, we find these observational studies sufficient for our purposes. First, our role is not to set a standard for the practice of medicine. Rather, we must simply confirm whether reasonable medical evidence supports the use of hydroxychloroquine as an early COVID-19 treatment, and we determine that a collection of large-scale observational studies suffices for that purpose. Second, a seminal review of the scientific literature has revealed that "on average, there is little evidence for significant effect estimate differences between observational studies and RCTs, regardless of specific observational study design, heterogeneity, or inclusion of studies of pharmacological interventions."²³⁵ There is thus no basis to cast aside the observational studies demonstrating hydroxychloroquine's efficacy as an early COVID-19 treatment.

233 Id.

Harvey A. Risch, Hydroxychloroquine in Early Treatment of High-Risk COVID-19 Outpatients: Efficacy and Safety Evidence, at 11 (Jun. 17, 2021), available at <u>https://earlycovidcare.org/wp-content/uploads/2021/09/Evidence-Brief-Risch-v6.pdf</u> (last visited Oct. 14, 2021).

Andrew Anglemyer et al., Healthcare outcomes assessed with observational study designs compared with those assessed in randomized trials, Cochrane Database of Systematic Reviews, at 1 (2014), available at <u>https://www.cochranelibrary.com/cdsr/doi/10.1002/14651858.MR000034.pub2/</u> epdf/full (last visited Oct. 14, 2021).

²³⁰ Million, supra, at 1070.

²³¹ Id.

Paul E. Alexander et al., Early multidrug treatment of SARS-CoV-2 infection (COVID-19) and reduced mortality among nursing home (or outpatient/ambulatory) residents, Medical Hypotheses, at 1 (2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8178530/pdf/main.pdf</u> (last visited Oct. 14, 2021).

We turn now to discuss the use of hydroxychloroquine as a prophylaxis, and although the data on that point seem to be smaller, there is some evidence suggesting that it might work for that purpose too. One study was a RCT of migrant workers quarantined in a large dormitory in Singapore, and it compared a group who used hydroxychloroquine as a prophylaxis to a group that received only vitamin C.236 The hydroxychloroquine group included 432 people, and only 31 of them (7.2%) contracted COVID-19 with acute respiratory symptoms.237 In contrast, 619 individuals were in the vitamin C group, and 69 of them (11.1%) developed COVID-19 with acute respiratory symptoms.²³⁸ Thus, the researchers concluded that prophylaxis with hydroxychloroquine is "superior to oral vitamin C in reducing SARS-CoV-2 infection."239 Additionally, an observational study of healthcare workers in Bulgaria found that out of 156 workers who used hydroxychloroquine as a prophylaxis, none of them presented with COVID-19 symptoms.240 By contrast, in the group of 48 workers who did not take hydroxychloroquine, three of them developed a symptomatic case of COVID-19.241 These results prompted the administrators at the Bulgarian Cardiac Institute to start a prophylactic strategy for their workers that "includes alternative months of [hydroxychloroguine] intake (200 mg daily) and months without therapy."242 In addition to these studies, there are a few others, some of which suggest marginal benefits, and some of which suggest that there might not be any. We are not aware of any of these studies showing serious adverse effects from use of low-dose hydroxychloroguine as a COVID-19 prophylaxis.

We pause here to reiterate that it is not our role to resolve the debate on hydroxychloroquine's effectiveness, either as an early COVID-19 treatment or as a preventative measure. These are matters for individual healthcare providers to assess based on the available data in consultation with their patients. Our only point is that reasonable data support the use of hydroxychloroquine as an early COVID-19 treatment and as a prophylaxis, and in light of that, we cannot find clear and convincing evidence

238 Id.

239 Id. at 314.

Raymond Chee Seong Seet et al., Positive impact of oral hydroxychloroquine and povidone-iodine throat spray for COVID-19 prophylaxis: An open-label randomized trial, 106 International Journal of Infectious Diseases 314, 314 (2021), available at <u>https://www.ijidonline.com/action/showPdf?pii=S1201-9712%2821%2900345-3</u> (last visited Oct. 14, 2021).

²³⁷ Id. at 319.

²⁴⁰ Iana Simova et al., Hydroxychloroquine for prophylaxis and treatment of COVID-19 in health-care workers, New Microbes and New Infections, at 1 (Nov. 2020), available at <u>https://www.sciencedirect.com/</u> <u>science/article/pii/S2052297520301657#</u>! (last visited Oct. 14, 2021).

²⁴¹ Id.

to file disciplinary actions against physicians who prescribe hydroxychloroquine for either of those purposes.

ii. Hydroxychloroquine, COVID-19, and Safety

During the pandemic, the FDA raised questions about hydroxychloroquine and adverse cardiac events.²⁴³ These kinds of concerns prompted one group of scholars to conduct a systematic review of the hydroxychloroquine safety literature pre-COVID-19. Their review of the data indicated that people taking that medication in appropriate doses "are at very low risk of experiencing cardiac [adverse events], particularly with short term administration" of the drug.²⁴⁴ The pre-COVID-19 data showed that heart issues occurred—albeit infrequently—only when patients took hydroxychloroquine in dangerously high doses or for many years on end.²⁴⁵

As to the increase of adverse cardiac events associated with COVID-19, the researchers questioned the prevalence of the problem by noting that several COVID-19 studies recorded "the use of [hydroxychloroquine] at variable doses without significant cardiac toxicity."²⁴⁶ They also observed that COVID-19 itself often causes heart issues. As they explained, "[t]he underlying pathophysiology of SARS-CoV-2 contributes to cardiac complications in the population it infects, with estimates ranging from 20-40% incidence."²⁴⁷ In particular, "[c]ardiac complications of cytokine storm have been well documented to involve fatal cardiac dysrhythmias and acute systolic heart failure."²⁴⁸ These researchers thus concluded that "the reported increased arrhythmic events in the COVID-19 era appear to be more related with the direct inflammatory effect of the virus (myocarditis) or the concomitant administration of multiple drugs capable of prolonging QT intervals rather than to hydroxychloroquine itself."²⁴⁹ They did not seem to think the medication itself had "change[d] after 70 years" of widespread use.²⁵⁰

- 247 Id. at 392.
- 248 Id. at 393.
- 249 Id. at 394.
- 250 Id.

²⁴³ U.S. Food and Drug Administration, FDA cautions against use of hydroxychloroquine or chloroquine for COVID-19 outside of the hospital setting or a clinical trial due to risk of heart rhythm problems, <u>https://www.fda.gov/drugs/drug-safety-and-availability/fda-cautions-against-use-hydroxychloro guine-or-chloroquine-covid-19-outside-hospital-setting-or</u> (last visited Oct. 14, 2021).

²⁴⁴ Fram, supra, at 391.

²⁴⁵ Id. at 390–92.

²⁴⁶ Id. at 393.

Others echoed these views. Another group reviewed the relevant studies and observed that "[m]ost of the available and credible data suggest that [hydroxychloroquine] is a safe drug."²⁵¹ That includes the pre-COVID-19 data—in "decades of . . . use by rheumatologists, . . . cardiac toxicity was rarely ever seen"—as well as the COVID-19-related studies—for example, the RECOVERY trial found "no cardiotoxicity" by hydroxychloroquine.²⁵² Indeed, the RECOVERY trial "prove[d] that [hydroxychloroquine] did not increase cardiac complications in COVID-19 cases despite using 4 times higher dosage than that used by rheumatologists."²⁵³ These authors also emphasized that "[m]ultiple mechanisms cause cardiac complications in patients with COVID-19 infection";²⁵⁴ thus, the infection's propensity to cause "intrinsic cardiac abnormalities . . . is probably acting as a confounder."²⁵⁵

Still another set of researchers reevaluated hydroxychloroquine's safety during the pandemic. They conducted a "meta-analysis to compare the safety of [hydroxychloroquine] versus placebo" for any indication.²⁵⁶ Although their "meta-analysis of RCTs found a significantly higher risk of skin pigmentation [issues] in [hydroxychloroquine] users versus placebo," they did not find any statistically significant increases in other adverse events, including "cardiac toxicity."²⁵⁷

In addition to these data tending to confirm hydroxychloroquine's safety when used in appropriate doses, a few other factors further lessen the cardiac concerns. For starters, one piece of key evidence contributing to the safety concerns surrounding hydroxychloroquine rested on admittedly fraudulent data. As discussed above, it was a study published in the Lancet on May 22, 2020.²⁵⁸ That study claimed that hydroxychloroquine was "associated with . . . an increased frequency of ventricular

257 Id.

258 Mehra, supra.

Shivraj Padiyar & Debashish Danda, Revisiting cardiac safety of hydroxychloroquine in rheumatological diseases during COVID-19 era: Facts and myths, 8 European Journal of Rheumatology 100, 100 (2021), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC8133889/pdf/ejr-8-2-100.pdf</u> (last visited Oct. 14, 2021).

²⁵² Id.

²⁵³ Id. at 102.

²⁵⁴ Id. at 102.

²⁵⁵ Id. at 100.

Khalid Eljaaly et al., Hydroxychloroquine safety: A meta-analysis of randomized controlled trials, Travel Medicine and Infectious Disease at 1 (Jul./Aug. 2020), available at <u>https://www.ncbi.nlm.nih.gov/pmc/articles/PMC7342171/</u> (last visited Oct. 14, 2021).

arrhythmias when used for treatment of COVID-19."²⁵⁹ That supposed finding was so startling that "major drug trials" involving hydroxychloroquine "were immediately halted";²⁶⁰ the WHO started pressuring countries like Indonesia that were widely using hydroxychloroquine to ban it;²⁶¹ and some countries—including France, Italy, and Belgium—decided to stop using it for COVID-19.²⁶²

The problem, however, is that the study was based on false data from a company named Surgisphere, whose founder and CEO Sapan Desai was a co-author on the published paper.²⁶³ The data were so obviously flawed that journalists and outside researchers began raising concerns within days of the paper's publication.²⁶⁴ Even the Lancet's editor in chief, Dr. Richard Horton, admitted that the paper was a "fabrication," "a monumental fraud, "²⁶⁵ and "a shocking example of research misconduct in the middle of a global health emergency."²⁶⁶ Approximately two weeks after its publication, the paper was retracted.²⁶⁷ An article published in *The Guardian* declared that "[g]iven the seriousness of the topic and the consequences of the paper, this [was] one of the most consequential retractions in modern history."²⁶⁸ Despite calls to "publish full explanations"

- 263 Boseley & Davey, supra.
- 264 Davey, supra.
- 265 Rabin, supra.
- 266 Boseley & Davey, supra.
- 267 Id.
- 268 Heathers, supra.

²⁵⁰ Id. at 1.

James Heathers, The Lancet has made one of the biggest retractions in modern history. How could this happen?, The Guardian (Jun. 5, 2020), available at https://www.theguardian.com/commentisfree/2020/jun/05/lancet-had-to-do-one-of-the-biggest-retractions-in-modern-history-how-could-this-happen (last visited Oct. 14, 2021).

Kate Lamb & Tom Allard, Indonesia, major advocate of hydroxychloroquine, told by WHO to stop using it, Reuters (May 26, 2020), available at <u>https://www.reuters.com/article/us-health-coronavirusindonesia-chloroqu/exclusive-indonesia-major-advocate-of-hydroxychloroquine-told-by-who-to-stopusing-it-idUSKBN23227L (last visited Oct. 14, 2021).</u>

²⁸² France, Italy, Belgium act to stop use of hydroxychloroquine for COVID-19 on safety fears, Reuters (May 27, 2020), available at <u>https://www.reuters.com/article/health-coronavirus-hydroxychloroquine-fr/update-1-france-italy-belgium-act-to-stop-use-of-hydroxychloroquine-for-covid-19-on-safety-fearsidUKL1N2D911J (last visited Oct. 14, 2021).</u>

of what happened," the Lancet has "declined to provide details regarding the retracted stud[y]."269

Further reducing the cardiac concerns is important information on the FDA's own website. The FDA "cautions against use of hydroxychloroquine . . . for COVID-19 *outside* of the hospital setting or a clinical trial due to risk of heart rhythm problems."²⁷⁰ But the agency's referenced support for this cautionary statement concerning *nonhospitalized* patients is its "review of safety issues with the use of hydroxychloroquine . . . to treat *hospitalized patients* with COVID-19."²⁷¹ It is questionable, however, to theorize about risks to nonhospitalized patients with mild COVID-19 based on data about heart issues in hospitalized patients with severe COVID-19 because, as explained above, cardiac complications often accompany the late stages of COVID-19. The FDA's concerns thus derive from a context—using hydroxychloroquine to treat hospitalized patients—that we are not addressing in this opinion.

It is important to note that although the medical literature tends to confirm that hydroxychloroquine is a safe medication when used in appropriate doses, any concerns about heart issues, even if resting on limited evidence, are serious. Prevailing principles of informed consent likely require physicians who present patients with the option of using hydroxychloroquine for early treatment of COVID-19 to inform them about the cardiac concerns that the FDA has identified. Also, for patients who have underlying cardiac issues, physicians should carefully consider whether hydroxychloroquine is the right choice for them. Finally, physicians should pay attention to which drugs they combine with hydroxychloroquine and evaluate the potential cardiac risks of those combinations. Failure to take such precautions could result in disciplinary action.

iii. U.S. Public Health Agencies on Hydroxychloroquine

The public health agencies in the United States have addressed the topic of hydroxychloroquine and COVID-19. The NIH "recommends against" its use "for the treatment of COVID-19 in hospitalized patients . . . and in nonhospitalized patients."²⁷² To justify its position against hydroxychloroquine for nonhospitalized patients, the NIH relied heavily on a RCT conducted by Mitja.²⁷³ While that study did not show great advantages in the hydroxychloroquine group, that group did have, as the NIH's own

²⁶⁹ Rabin, supra.

²⁷⁰ U.S. Food and Drug Administration, FDA cautions against use of hydroxychioroquine or chloroquine for COVID-19 outside of the hospital setting or a clinical trial due to risk of heart rhythm problems, <u>https://www.fda.gov/drugs/drug-safety-and-availability/fda-cautions-against-use-hydroxychloroguine-or-chloroquine-covid-19-outside-hospital-setting-or</u> (last visited Oct. 14, 2021) (emphasis added).

²⁷¹ Id. (emphasis added).

²⁷² NIH, COVID-19 and Hydroxychloroquine, supra.

website reports, a slight reduction in the risk of hospitalization (7.1% risk in the control arm versus 5.9% risk in the treatment arm) and in the time to resolution of symptoms (12 days in the control arm versus 10 days in the treatment arm).²⁷⁴ As for serious adverse events, more (12) were reported in the control group than the hydroxychloroquine group (8), and the researchers determined that the serious adverse events in the hydroxychloroquine group were not related to the drug.²⁷⁵ Thus, this study, particularly when considered in light of the large-scale observational studies discussed above, appears to be an insufficient basis to definitively recommend against using hydroxychloroquine as an early COVID-19 treatment.

The FDA, for its part, has questioned not only hydroxychloroquine's safety, as we discussed above, but also its efficacy. The agency's position grew out of its approval and subsequent disapproval of an Emergency Use Authorization (EUA) involving hydroxychloroquine. That EUA was issued on March 28, 2020, and it authorized licensed healthcare providers to use hydroxychloroquine donated to the Strategic National Stockpile to treat patients hospitalized with COVID-19.²⁷⁶ Though this EUA was necessary to authorize the use of a specific source of hydroxychloroquine for a specific purpose, it was not required to allow healthcare providers to prescribe hydroxychloroquine off-label for COVID-19. That option was already available, as our prior discussion of off-label use makes clear. When the FDA revoked the EUA a few months later, on June 15, 2020, that is when it stated its current position on hydroxychloroquine and COVID-19.²⁷⁷

In that revocation, the FDA said that it no longer "believe[s] that oral formulations of [hydroxychloroquine]...may be effective in treating COVID-19" or that "that the known and potential benefits of these products outweigh their known and potential risks."²⁷⁸

Letter from Denise M. Hinton, Chief Scientist, U.S. Food and Drug Administration, to Dr. Rick Bright, Director of Biomedical Advanced Research and Development Authority (BARDA), Office of Assistant Secretary for Preparedness and Response (ASPR), U.S. Department of Health and Human Services (HHS) (Mar. 28, 2020), available at https://www.fda.gov/media/136534/download (last visited Oct. 14, 2021).

²⁷⁷ Letter from Denise M. Hinton, Chief Scientist, U.S. Food and Drug Administration, to Gary L. Disbrow, Deputy Assistant Secretary, Director of Medical Countermeasure Programs, Biomedical Advanced Research and Development Authority (BARDA), Office of Assistant Secretary for Preparedness and Response (ASPR), U.S. Department of Health and Human Services (HHS) (Jun. 15, 2020), available at https://www.fda.gov/media/138945/download (last visited Oct. 14, 2021).

278 Id. at 2.

National Institutes of Health, Table 2b. Chloroquine or Hydroxychloroquine and/or Azithromycin: Selected Clinical Data, <u>https://www.covid19treatmentguidelines.nih.gov/tables/table-2b/</u> (last visited Oct. 14, 2021) (discussing Oriol Mitjà, *Hydroxychloroquine for Early Treatment of Adults With Mild Coronavirus Disease 2019: A Randomized, Controlled Trial*, Clinical Infectious Diseases (2020), *available at* <u>https://academic.oup.com/cid/advance-article/doi/10.1093/cid/ciaa1009/5872589</u> (last visited Oct. 14, 2021)).

²⁷⁵ Id. (discussing Mitjà, supra).

Because both the EUA and its revocation deal only with hydroxychloroquine's use in hospitalized patients, they do not address the treatment topic that we are considering in this opinion—hydroxychloroquine's use as an early COVID-19 treatment.

The FDA's EUA revocation included four justifications, none of which establishes let alone by clear and convincing evidence—that hydroxychloroquine is ineffective as an early treatment of COVID-19. First, the FDA said that the "suggested dosing regimens... are unlikely to produce an antiviral effect" because they will not create sufficient "drug concentration" in the body.²⁷⁹ But as the FDA's revocation itself acknowledged, hydroxychloroquine's "immunomodulatory effects," as opposed to its antiviral effects, are not "predicated on achieving [certain hydroxychloroquine] concentration[]" levels.²⁸⁰ Moreover, the FDA based its views on the assumption that "free drug concentration in the plasma" are "likely to be equal to free extracellular tissue concentration."²⁸¹ But other researchers' simulations showed that hydroxychloroquine's "concentration in lung tissue was much higher than in plasma,"²⁸² leading them to conclude that moderate doses are "recommended to treat SARS-CoV-2 infection."²⁸³ Thus, the FDA's pessimism about hydroxychloroquine's potential antiviral capacity is open to reasonable debate in the scientific community.

Second, the FDA wrote that "[e]arlier reports of decreased viral shedding" with hydroxychloroquine "treatment have not been consistently replicated."²⁸⁴ Notice that the FDA did not say that the studies have *disproven* a reduction in viral shedding; rather, the agency recognized that the evidence was still evolving and that some studies did in fact observe a positive "impact on viral shedding."²⁸⁵ This criticism, on its face, is thus insufficient to dismiss hydroxychloroquine's use as an early COVID-19 intervention. Additionally, doubts about hydroxychloroquine's effect on viral shedding question only one of the drug's many possible mechanisms of action against COVID-19. More salient

²⁷⁹ U.S. Food and Drug Administration, Memorandum Explaining Basis for Revocation of Emergency Use Authorization for Emergency Use of Chloroquine Phosphate and Hydroxychloroquine Sulfate, at 1, 4, available at <u>https://www.fda.gov/media/138945/download</u> (last visited Oct. 14, 2021) (hereinafter, "FDA EUA Revocation Memo").

²⁸⁰ Id. at 4.

²⁸¹ Id.

Xueting Yao et al., In Vitro Antiviral Activity and Projection of Optimized Dosing Design of Hydroxychloroquine for the Treatment of Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), Clinical Infectious Diseases, at 13 (2020), available at <u>https://www.ncbi.nlm.nih.gov</u> /pmc/articles/PMC7108130/pdf/ciaa237.pdf (last visited Oct. 14, 2021).

²⁸³ Id. at 2.

²⁸⁴ FDA EUA Revocation Memo, supra, at 1.

²⁸⁵ Id. at 6.

information is whether the drug is actually cecreasing hospitalization and mortality rates when used as an outpatient treatment. As we discussed above, many large observational studies strongly suggest that hydroxychloroquine does in fact keep people diagnosed with COVID-19 out of the hospital and alive. That evidence is far more relevant of the drug's potential efficacy as an early COVID-19 treatment than debates about viral shedding.

Third, the FDA found it compelling that "NIH guidelines now recommend against" using hydroxychloroquine "outside of a clinical trial."²⁸⁶ But as previously explained, the NIH's recommendation concerning COVID-19 outpatients does not rest on undisputed support. Thus, the NIH's guidelines should not be considered a basis upon which to ban healthcare providers from using hydroxychloroquine for COVID-19.

Fourth, the FDA stressed that "[r]ecent data from a large randomized controlled trial"—the RECOVERY trial mentioned above—"showed no evidence of benefit . . . of [hydroxychloroquine] treatment in hospitalized patients with COVID-19."²⁸⁷ Yet as we have already discussed, a study about hospitalized patients does not address hydroxychloroquine's efficacy as an outpatient COVID-19 treatment. Indeed, the RECOVERY team itself reported that while its "findings indicate that hydroxychloroquine is not an effective treatment for hospitalized patients with Covid-19," it does "not address [the drug's] use as prophylaxis or in patients with less severe SARS-CoV-2 infection managed in the community."²⁸⁸ In sum, none of the FDA's four reasons, in isolation or taken together, clearly establish that hydroxychloroquine is ineffective as an early treatment against COVID-19.

Despite raising doubts about hydroxychloroquine's use against COVID-19, the FDA has consistently affirmed that healthcare providers retain the right to use hydroxychloroquine as a part of early COVID-19 treatment. At least four statements demonstrate this.

First, the FDA's current website says (and has said since July 2020) that "[i]f a healthcare professional is considering use of hydroxychloroquine or chloroquine to treat or prevent COVID-19, FDA recommends checking www.clinicaltrials.gov for a suitable clinical trial and consider enrolling the patient." This plainly assumes that healthcare providers have the right to use hydroxychloroquine to treat COVID-19.

Second, on May 29, 2020, then-FDA Commissioner Stephen Hahn acknowledged that "[m]any physicians have . . . prescribed [hydroxychloroquine] for patients with COVID-19 based on an individual assessment of the potential benefits versus the risks

²⁸⁶ *Id.* at 1.

²⁸⁷ Id.

RECOVERY Collaborative Group, Effect of Hydroxychloroquine in Hospitalized Patients with Covid-19, 383 The New England Journal of Medicine 2030, 2038 (Nov. 2020), available at https://www.nejm.org/doi/pdf/10.1056/NEJMoa2022926?articleTools=true (last visited Oct. 14, 2021).

for an individual patient."²⁸⁹ He added that "[p]rescribing a product for uses not specifically included in the official labeling is common in the practice of medicine" and that the FDA does not "prohibit[] physicians from prescribing medications" because the agency does "not regulate the practice of medicine."²⁹⁰ These statements are still posted on the FDA's website, and we are not aware of any subsequent FDA statements revoking them.

Third, in June 2020, after the FDA revoked the hydroxychloroquine EUA, Health and Human Services Secretary Alex Azar said: "At this point, hydroxychloroquine and chloroquine are just like any other approved drug in the United States. They may be used in hospital, they may be used in out-patient, they may be used at home—all subject to a doctor's prescription."²⁹¹ Leaving no doubt about this point, Secretary Azar added that "[i]f a doctor wishes to prescribe [hydroxychloroquine], working with a patient, they may prescribe it for any purpose that they wish."²⁹² We are not aware of any subsequent statement revoking this guidance.

Fourth, in late July 2020, then-FDA Commissioner Hahn reiterated that "whether people should take hydroxychloroquine as a treatment" for COVID-19 is a decision that "should be made between a doctor and a patient."²⁹³ He specifically stated: "A doctor and a patient need to assess the data that's out there, FDA does not regulate the practice of medicine, and that in the privacy of the doctor-patient relationship is where that decision should be made."²⁹⁴

iv. Foreign Public Health Agencies, Professional Associations, and Physicians on Hydroxychloroquine

The WHO "recommend[s] against administering hydroxychloroquine . . . for treatment of COVID-19" for "patients with any disease severity and any duration of symptoms." ²⁹⁵ It reached this recommendation after concluding that hydroxychloroquine

292 Id.

²⁹³ Tal Axelrod, FDA chief: Hydroxychloroquine use a decision between doctor and patient, The Hill (Jul. 30, 2020), <u>https://thehill.com/policy/healthcare/509733-fda-chief-hydroxychloroquine-use-a-decisionbetween-doctor-and-patient?rl=1</u> (last visited Oct. 14, 2021).

²⁸⁹ FDA, Bringing Perspective, supra.

²⁹⁰ Id.

²⁹¹ Trump White House Archives, Remarks by President Trump in Roundtable Discussion on Fighting for America's Seniors (Jun. 15, 2020), available at <u>https://trumpwhitehouse.archives.gov/briefingsstatements/remarks-president-trump-roundtable-discussion-fighting-americas-seniors/</u> (last visited Oct. 14, 2021).

²⁹⁵ WHO COVID-19 Guidelines, supra, at 26.

"probably do[es] not reduce mortality" and that its "effect on . . . admission to hospital . . . remains uncertain."²⁹⁶ To the extent that this recommendation purports to address hydroxychloroquine's effectiveness as an early treatment for COVID-19, it arguably rests on weak evidence. Although it is difficult to determine how many of the studied individuals were outpatients, it appears that most were hospitalized. For instance, the WHO says that it consulted 29 studies in concluding that "[h]ydroxychloroquine probably does not reduce mortality," but the only study specifically cited is the RECOVERY trial,²⁹⁷ which, as we already indicated, included only patients hospitalized with COVID-19.²⁹⁸ In addition, the WHO's statistics on hospitalization rates, which consisted of one RCT that included 465 outpatients, suggests hydroxychloroquine's efficacy.²⁹⁹ That trial revealed a hospitalization rate of 47 per 1,000 people in the control group but only 19 of 1,000 people in the hydroxychloroquine arm.³⁰⁰ It thus seems as if the WHO may have overreached in definitively declaring that hydroxychloroquine holds no promise as an early COVID-19 treatment.

The WHO also "recommend[s] against administering hydroxychloroquine prophylaxis to individuals who do not have COVID-19" because it believes that prophylaxis "hydroxychloroquine has a small or no effect on death and hospital admission" and that it "probably has a small or no effect on laboratory-confirmed COVID-19."³⁰¹ Disagreeing with this, the team of researchers conducting the COPCOV trial on prophylaxis hydroxychloroquine has announced that the WHO's conclusions are "scientifically unsound."³⁰² In their statement on this topic, the COPCOV team explained that the available RCTs "suggest substantial uncertainty as to the benefit of hydroxychloroquine in preventing COVID-19," but the "overall trend [is] towards benefit."³⁰³

298 RECOVERY Collaborative Group, supra, at 2030.

- 299 WHO COVID-19 Guidelines, supra, at 29.
- 300 Id.

World Health Organization, WHO Living guideline: Drugs to prevent COVID-19, at 12 (Mar. 2, 2021), available at <u>https://apps.who.int/iris/bitstream/handle/10665/339877/WHO-2019-nCoV-prophylaxes-2021.1-eng.pdf?sequence=13&isAllowed=y</u> (last visited Oct. 14, 2021).

³⁰² The COPCOV Trial's position statement on "A living WHO guideline on drugs to prevent COVID-19," MORU Tropical Health Network (Mar. 5, 2021), <u>https://www.tropmedres.ac/news/copcov-response-tolatest-who-guidelines-on-hydroxychloroquine-for-covid-19-trials-1</u> (last visited Oct. 14, 2021).

²⁹⁶ Id. at 27.

²⁹⁷ *Id.* at 28.

As for the professional associations' and physician groups' views on hydroxychloroquine, it appears that they generally adopt the same position they took on ivermectin. Those like the AAPS that support ivermectin as an option for early COVID-19 treatment generally support hydroxychloroquine too, while those like the AMA, APhA, and ASHP that oppose one typically resist the other. Additionally, many physician groups use early COVID-19 treatment protocols that include hydroxychloroquine. For example, an article co-authored by over 50 doctors in Reviews in Cardiovascular Medicine outlines an early treatment protocol that includes hydroxychloroquine as a key component.³⁰⁴

Considering the evidence discussed above, we do not find that clear and convincing evidence would warrant disciplining physicians who prescribe hydroxychloroquine for the prevention or early treatment of COVID-19 after first obtaining informed patient consent.

CONCLUSION

Based on the available data, we do not find clear and convincing evidence that a physician who first obtains informed consent and then utilizes ivermectin or hydroxychloroquine for COVID-19 violates the UCA. This conclusion is subject to the limits noted throughout this opinion. Foremost among them are that if physicians who prescribe ivermectin or hydroxychloroquine neglect to obtain informed consent, deceive their patients, prescribe excessively high doses, fail to check for contraindications, or engage in other misconduct, they might be subject to discipline, no less than they would be in any other context.

As we have stressed throughout, this opinion is based only on the data and information available at this time. If the relevant medical evidence materially changes, that could impact our conclusions. Also, though an opinion from our office about possible UCA violations would ordinarily focus on healthcare practices within Nebraska, the context of a global pandemic necessitates looking for evidence far beyond our State's borders, as we have done here. Thus, the analytical roadmap in this opinion likely has limited application outside the circumstance of a global pandemic.

We emphasize in closing that our office is not recommending any specific treatments for COVID-19. That is not our role. There are multiple treatment options outside the scope of this opinion—including treatments that have been officially approved by the FDA—that physicians and their patients should carefully consider. This opinion takes no position on them. Rather, we address only the off-label early treatment options discussed in this opinion and conclude that the available evidence suggests that they might work for some people. Allowing physicians to consider these early treatments will free them to evaluate additional tools that could save lives, keep patients out of the hospital, and provide relief for our already strained healthcare system.

³⁰⁴ McCullough, Multifaceted, supra, at 522-23.

Very truly yours,

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