Will the antimalarial drug take over to combat COVID-19?

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Abstract

Background China has been fighting the epidemic of pneumonia-like diseases first detected for over a month in the city of Wuhan in December 2019. The disease epidemic is caused by a novel coronavirus, called COVID-19, which has now infected more than 700,000 people worldwide. With a death toll approaching that of China’s SARS-CoV outbreak in 2002 and 2003, 2019-nCoV has contributed to an international emergency in public health, placing all health organizations on high alert. Such large numbers of infected and deceased people require an urgent need for reliable, inexpensive, and cheap drugs to control and reduce the outbreak.

Objective To systematically review and evaluate the pattern of COVID-19 and the treatment plans.

Methods This systematic review was conducted according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines. The articles were searched from databases like PubMed, the Cochrane Library, ScienceDirect, and the Health Research and Development Information Network (HERDIN) combining MeSH and free-text terms.

Results This analysis highlights the agent of COVID-19 and the possible transmission. The current research taking place to overcome this complex disease and the urgent need to develop improved therapeutics are also discussed.

Conclusion Herein, we present an epidemiological overview of the currently available information on the treatment claimed to have helped to bring the situation under control.

Keywords Wuhan · China · COVID-19 · Outbreak · Chloroquine · Hydroxychloroquine

Introduction

Several cases of unknown etiology pneumonia have been confirmed and registered in Wuhan, Hubei Province, China since December 8, 2019 (Huang et al. 2020). Most of the sufferers worked or lived around the local Huanan Seafood Wholesale Market, where live animals were on sale. Extreme acute respiratory infection signs occurred in the early stages of this pneumonia, with some sufferers quickly developing acute respiratory distress syndrome (ARDS), severe breathing failure, and other significant complications. On January 7, 2020, a single coronavirus was detected from a patient’s throat swab sample by the Chinese Center for Disease Control and Prevention (China CDC) and was eventually called COVID-19 by the World Health Organization (WHO) (Ji et al. 2020). This virus has now caused a total of 750,890 confirmed human infections worldwide, with 36,405 deaths (WHO Situation report - 71, 31 March 2020 2020). The whole world looks forward to some kind of relief and treatment for this serious pandemic.

Agent

Coronaviruses belong to the Coronavirinae subfamily of the Coronaviridae family of the order Nidovirales, and this subfamily consists of four genera: Alphacoronavirus, Betacoronavirus, Gammaporonavirus, and Deltacoronavirus (Chen et al. 2020). Coronaviruses of zoonotic origin can, however, evolve into a strain capable of infecting mortally ill people (Su et al. 2016).

COVID-19 is the seventh MERS-nCoV and SARS-nCoV coronavirus family member to infect humans. A genetic sequence of the virus was made available to the WHO via genome sequencing, which allowed various laboratories to
produce diagnostic reverse transcription polymerase chain reaction (RT-PCR) tests specifically for viral RNA detection. COVID-19 is a category 2B β CoV with a genetic sequence similarity of more than 70% to SARS-nCoV (Hui et al. 2020). The origins of the new infection with coronavirus was ascertained as bats. With full genome sequences, Zhou and colleagues found that the virus is 96% the same as a bat coronavirus at the level of the entire genome (Zhou et al. 2020). Wu and colleagues performed a phylogenetic analysis of the entire viral genome and deduced that the virus was most closely associated with a group of previously isolated SARS-like coronaviruses from bats in China (Wu et al. 2020).

### Possible transmissions

After the first cases of the COVID-19 disease were linked to direct access to Wuhan’s Huanan Seafood Wholesale Market, it was suspected that the main mechanism was a transmission from animal to human. The cycle of adaptation was not associated with subsequent incidents, however. It was also known that the virus could also be transmitted from human to human and that the most frequent cause of COVID-19 spread is symptomatic individuals. The probability of transmission appears unlikely before symptoms arise, but it cannot be ruled out. There are also indications that the virus can be transmitted by people who remain asymptomatic.

Based on data from the first cases in Wuhan and inquiries performed by the Chinese CDC and local CDCs, the incubation period could typically be between 3 to 7 days and up to 2 weeks, as the longest duration from infection to symptoms was 12.5 days [95% confidence interval (CI), 9.2 to 18] (Li et al. 2020).

These data also showed that this novel outbreak doubled every 7 days, although the simple number of reproductions ($R_0$) was 2.2. In other words, every patient transmits the infection to an additional 2.2 individuals on average. Of note, estimates for the SARS-CoV outbreak $R_0$ were about 3 in 2002–2003 (Bau et al. 2005).

### Treatment approaches so far

COVID-19 is not approved for specific antiviral therapy and does not have a vaccine. The condition is symptomatic, and the main treatment intervention for severely compromised patients is oxygen therapy. Mechanical ventilation may be needed in cases of respiratory failure refractory to oxygen therapy, while hemodynamic support is necessary for septic shock control.

On January 28, 2020, the WHO issued a paper summarizing guidelines from the WHO and clinical evidence collected from the treatment of previous HCoV epidemics. This study addresses measures to classify and sort patients with extreme acute respiratory disease, methods to avoid and contain infections, early intervention and monitoring therapy, guidelines for laboratory diagnosis, management of respiratory failure and ARDS, management of septic shock, prevention of complications, treatments, and consequences for pregnant patients (Cascella et al. 2020).

### The launch of a megatrial

A combination of the drugs used against HIV, a malaria vaccine first tested during the Second World War, and a new antiviral against Ebola last year whose promise waned. On March 20, 2020, the WHO initiated a large global project, called SOLIDARITY, to find out if any of these drugs can be used with the new coronavirus infections to cure the deadly respiratory illness. The project also examines unapproved medications that have performed well with the other two deadly coronaviruses in animal studies, causing severe acute respiratory syndrome (SARS) and Middle East respiratory syndrome (MERS) (Kupferschmidt and Cohen 2020).

### Will the approved or investigational drug take over?

There are no medications specifically licensed by the US Food and Drug Administration (FDA) for the treatment of COVID-19 patients. In several hundred clinical trials that are ongoing across the globe, many drugs approved for other indications as well as other investigational drugs are being studied. Two of the approved drugs (chloroquine and hydroxychloroquine) and one of the commonly used research agents (remdesivir) are in the limelight for the whole world on that basis (Centers for Disease Control and Prevention, CDC 2020).

### Remdesivir

Remdesivir is a large-scale intravenous antiviral investigative drug that inhibits viral replication by premature termination of RNA transcription and has in vitro activity against SARS-CoV-2 and in vitro and in vivo activity against related beta coronaviruses (Mulangu et al. 2019).

Recently, remdesivir was recognized as a promising antiviral drug against a large array of RNA viruses. This is equivalent to adenosine, which integrates into the chains of embryonic viral RNA and results in premature termination. Warren et al. demonstrated that the non-human primate model’s intravenous administration of 10 mg/kg of remdesivir resulted in concomitant persistent levels of its active form in the blood.
COVID-19 was 6.90 μM in lung. The EC90 value of chloroquine in Vero E6 cells against SARS-CoV cell receptor glycosylation (Vincent et al. 2005). In addition to its antiviral function, chloroquine has an immune-modulating role that can improve synergistically its antiviral effect in vivo. Upon oral administration, chloroquine is spread extensively throughout the entire body, including the lung. The EC90 value of chloroquine in Vero E6 cells against COVID-19 was 6.90 μM, which can be clinically attainable as seen in the plasma of patients receiving 500 mg of rheumatoid arthritis. The time-of-addition assay demonstrated, in an in vitro analysis, that chloroquine performed at both entry and post-entry stages of COVID-19 infections in Vero E6 cells (Wang et al. 2020).

**Chloroquine**

Chloroquine, a widely used drug for antimalarial and autoimmune disorders, has been recently described as a possible broad-spectrum antiviral medicine. Chloroquine is known to prevent viral infection by growing the endosomal pH needed for virus/cell fusion, as well as interfering with SARS-CoV cell receptor glycosylation (Vincent et al. 2005). In addition to its antiviral function, chloroquine has an immune-modulating role that can improve synergistically its antiviral effect in vivo. Upon oral administration, chloroquine is spread extensively throughout the entire body, including the lung. The EC90 value of chloroquine in Vero E6 cells against COVID-19 was 6.90 μM, which can be clinically attainable as seen in the plasma of patients receiving 500 mg of rheumatoid arthritis. The time-of-addition assay demonstrated, in an in vitro analysis, that chloroquine performed at both entry and post-entry stages of COVID-19 infections in Vero E6 cells (Wang et al. 2020).

**Hydroxychloroquine**

Hydroxychloroquine and chloroquine are cellular autophagy modulators that interact with enveloped viruses, such as retroviruses, flaviviruses, and coronaviruses, at late stages of replication (Savarino et al. 2003; Byrd and Horwitz 1991). Hydroxychloroquine or chloroquine anti-HIV-1 activity was demonstrated in conjunction with antiretroviral drugs such as zidovudine, hydroxyurea, and didanosine in vitro, or patients (Paton and Aboulhab 2005). Hydroxychloroquine raises the pH in intracellular vacuoles and improves processes such as protein degradation by acidic lysosome hydrolases, assembly of macromolecules in endosomes, and post-translation protein modification in the Golgi apparatus. It is suggested that the antirheumatic properties of these compounds derive from their involvement in macrophages and other antigen-presenting cells with the “antigen processing”. These compounds give greater confidence in aspects of the clinical and laboratory parameters. These are distinguished from glucocorticoids and non-steroidal anti-inflammatory agents by their slow start of the action.

**Conclusion**

This latest virus epidemic has challenged most countries across the globe with economic, medical, and public health infrastructures. Time alone can tell how the virus is going to affect our lives. Potential outbreaks of the viruses and pathogens of zoonotic origin are likely to occur. Hydroxychloroquine has been used for 70 years as a cheap and safe drug, and is, thus, potentially clinically important against COVID-19. There is hope in this high-need scenario that chloroquine could change the whole history of this pandemic assault.

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