Abstract

The Coronavirus disease 19 (COVID-19) is a highly transmittable and pathogenic viral infection caused by severe acute respiratory syndrome Coronavirus 2 (SARS-CoV-2), which emerged in Wuhan, China in December 2019. Since the emergence of the 2019 novel coronavirus (2019-nCoV), the World Health Organization (WHO) has declared it a public health emergency of international concern. A genomic study revealed that SARS-CoV-2 is phylogenetically related to severe acute respiratory syndrome (SARS) bat viruses; therefore, bats could be the possible primary reservoir. The intermediate source of origin and transfer to humans is not known, however, the rapid human-to-human transfer has been confirmed widely via droplets or direct contact, and infection has been estimated to have an incubation period of 6.4 days. Currently, controlling infection to prevent the spread of SARS-CoV-2 is the primary intervention being used. However, public health authorities should keep monitoring the situation closely, as the more we can learn about this novel virus and its associated outbreak, the better we can respond.

Keywords: SARS-CoV-2, COVID-19, China, Phylogenetic, Genomic analysis, Remdesivir

Résumé

Le COVID-19 est une nouvelle maladie mortelle avec des informations limitées sur sa transmissibilité, la gravité de ses séquelles, ses manifestations cliniques et l'épidémiologie. Ce commentaire a analysé l'épidémiologie mondiale du COVID-19 parmi la population vulnérable. L'analyse a révélé que la plupart des cas pédiatriques de COVID-19 ne sont pas graves, mais que des maladies graves associées surviennent toujours chez les enfants. Tous les âges des enfants sont sensibles au COVID-19 et aucune différence significative entre les sexes n'existe. L'infection au COVID-19 pendant la grossesse a eu des conséquences fatales pour les mères, mais moins mortelles pour le bébé. Les groupes de points chauds pour le COVID-19 sont les prisons / prisons, les maisons de soins infirmiers / de groupe et les établissements à long terme où résident la plupart des populations vulnérables. Les groupes ethniques minoritaires aux États-Unis et au Royaume-Uni sont exposés de manière disproportionnée à l'infection au COVID-19 et à la mort que les Caucasiens. La différence peut être due au fait que les minorités ethniques sont exposées à des risques plus élevés au travail et aux disparités structurelles économiques et sanitaires de longue date dans les deux pays. Il y a maintenant des changements dans les lignes directrices sur les personnes qualifiées pour recevoir des ventilateurs dans des situations difficiles dans de nombreux pays du monde si le système de santé est débordé.

Mots-clés: COVID-19, épidémiologie, population vulnérable

Introduction

Since the emergence of the 2019 novel coronavirus (2019-nCoV) in Wuhan, China, in December 2019, it has rapidly spread across China and many other countries. So far, 2019-nCoV has affected more than 5.5 million persons in more than 200 countries, making it a major global health concern. On 11th February 2020, the World Health Organization (WHO) announced a new name for the epidemic disease caused by 2019-nCoV: Coronavirus disease (COVID-19). Regarding the virus itself, the International Committee on Taxonomy of Viruses has renamed the previously provisionally named 2019-nCoV as severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2). Although early studies reported a link between a single local fish,
wild animal market and most cases of infection, indicating possible animal-to-human transmission, and studies have increasingly demonstrated human-to-human transmission of SARS-CoV-2 through droplets or direct contact. Moreover, according to one study, presumed hospital-related transmission of SARS-CoV-2 was suspected in 41% of patients. Based on the evidence of a rapidly increasing incidence of infections and the possibility of transmission by asymptomatic carriers, SARS-CoV-2 can be transmitted effectively among humans and exhibits high potential for a pandemic. In addition to the high transmission efficiency of SARS-CoV-2, the advancement and convenience of global travel could further enhance its worldwide spread. On 30th January 2020, the WHO declared the COVID-19 outbreak as the sixth public health emergency of international concern, following H1N1 (2009), polio (2014), Ebola in West Africa (2014), Zika (2016) and Ebola in the Democratic Republic of Congo (2019). Therefore, health workers, governments and the public needs to co-operate globally to prevent its spread. 

**Evolution**

Coronaviruses belong to the Coronaviridae family in the Nidovirales order. Coronaviruses are minute in size (65-125 nm in diameter) and contain a single-stranded RNA as a nucleic material, size ranging from 26 to 32 kbs in length (Figure 1). The subgroups of Coronaviruses are alpha, beta, gamma and delta Coronaviruses. Each of the subgroups of corona viruses has many serotypes. Some of them were affect human while other affected animals such as pigs, birds, cats, mice and dogs. 

In addition to seasonal influenza, reported pathogens of pneumonia include adenovirus, Coronavirus 229E/NL63/OC43, human bocavirus, human metapneumovirus, parainfluenza virus 1/2/3, rhinovirus and respiratory syncytial virus A/B. Moreover, these viruses can cause co-infection in the setting of community-acquired bacterial pneumonia. Using molecular methods, knowledge about the role of these viruses in the setting of pneumonia has achieved significant advancements. SARS-CoV-2 was found to be a positive-sense, single-stranded RNA virus belonging to the genus Betacoronavirus. Phylogenetic analysis revealed that SARS-CoV-2 is closely related (88-89% similarity) to two bat-derived SARS-like Coronaviruses, namely bat-SL-CoVZC45 (GenBank accession no. MG772933.1) and bat-SL-CoVZXC21 (GenBank accession no. MG772934.1), but it is more distant from SARS-CoV (~79% similarity) and Middle East respiratory syndrome Coronavirus (MERS-CoV) (~50% similarity). Chen et al. applied an RNA-based metagenomic next-generation sequencing approach to identify a human Coronavirus from two pneumonia cases during the Wuhan outbreak in 2019. Its entire genome was 29881 bp in length. Phylogenetic analysis indicates that SARS-CoV-2 is similar to the Coronavirus circulating in Rhinolophus (bats), with 98.7% nucleotide similarity to the partial RNA-dependent RNA polymerase (RdRp) gene of the bat Coronavirus strain BtCoV/4991 (GenBank KP876546, 370 bp sequence of RdRp) and 87.9% nucleotide similarity to bat Coronavirus strain bat-SL-CoVZC45 and bat-SL-CoVZXC21. Evolutionary analysis based on ORF1a/1b, S and N genes suggests that SARS-CoV-2 is more likely a novel Coronavirus that was independently introduced from animals to humans. Based on the findings of genomic investigations and the presence of some bats and live animals in the seafood market in Wuhan, SARS-CoV-2 may have originated from bats or bat droppings associated with contaminated materials in the market or surrounding region.

**Comparative analysis of emergence and spread of Coronaviruses**

First case of corona virus was notified as cold in 1960. According to the Canadian study 2001, approximately 500 patients were identified as Flu-like system. 17-18 cases of them were confirmed as infected with corona virus strain by polymerase chain reaction. Corona was treated as simple non fatal virus till 2002. In 2003, various reports published with the proofs of spreading the corona to many countries such as United States America, Hong Kong, Singapore, Thailand, Vietnam and in...
Taiwan. Several cases of severe acute respiratory syndrome caused by corona virus and then mortality more than 1000 patients was reported in 2003. This was the black year for scientist. After a deep exercise they conclude and understand the pathogenesis of this disease and discovered it was corona virus. But still total of 8096 patients were confirmed as infected with corona virus. So, in 2004, World health organization and centers for disease control and prevention declared a state emergency. Another study report of Hong Kong confirmed 50 patients of severe acute respiratory syndrome while 30 of them were confirmed as corona virus infected while in 2012, Saudi Arabian reported several infected patient and deaths\textsuperscript{34-36}. COVID-19 was first identified and isolated from pneumonia patient belongs to Wuhan, China.

**Epidemiology**

As of 25 May 2020, situation report from the WHO showed a total of 5,520,743 confirmed cases of COVID-19 globally\textsuperscript{36}. Since the outbreak of COVID-19 originated from Wuhan City, Hubei province, in China, there has been a steady increase in the daily total number of COVID-19 cases globally, both within and outside China. Regarding new cases of COVID-19, a declining trend was found globally, in China but not outside China, mainly in international conveyance (Japan) on 11th February 2020. Twenty-eight countries/regions have reported confirmed cases, including mainland China, Japan, Singapore, Hong Kong Special Administrative Region (SAR), Thailand, South Korea, Taiwan, Australia, Malaysia, Germany, Vietnam, the USA, Macao.
Figure 3: The key reservoirs and mode of transmission of Coronaviruses (suspected reservoirs of SARS-CoV-2 are red encircled); only a and b Coronaviruses have the ability to infect humans, the consumption of infected animal as a source of food is the major cause of animal to human transmission of the virus and due to close contact with an infected person, the virus is further transmitted to healthy persons. Dotted black arrow shows the possibility of viral transfer from bat whereas the solid black arrow represents the confirmed transfer.

Figure 4: The life cycle of SARS-CoV-2 in host cells; begins its life cycle when S protein binds to the cellular receptor ACE2. After receptor binding, the conformation change in the S protein facilitates viral envelope fusion with the cell membrane through the endosomal pathway. Then SARS-CoV-2 releases RNA into the host cell. Genome RNA is translated into viral replicase polyproteins pp1a and 1ab, which are then cleaved into small products by viral proteinases. The polymerase produces a series of sub genomic mRNAs by discontinuous transcription and finally translated into relevant viral proteins. Viral proteins and genome RNA are subsequently assembled into virions in the ER and Golgi and then transported via vesicles and released out of the cell. ACE2, angiotensin-converting enzyme 2; ER, endoplasmic reticulum; ERGIC, ER–Golgi intermediate compartment.
SAR, the United Arab Emirates, Canada, France, the Philippines, the UK, Italy, India, Russia, Finland, Sweden, Sri Lanka, Cambodia, Nepal, Spain and Belgium. China has had the largest number of patients with COVID-19 (n= 42690), followed by Singapore (n= 45). Asia has had most of the reported cases, followed by Europe, North America and Australia, but no cases have been reported in Africa. Within China, Hubei has endured the largest number of infected patients (n = 31,728), followed by Guangdong (n = 1177), Zhejiang (n = 1117) and Henan (n = 1105). A total of 1017 mortalities have been reported globally with only 2 mortalities occurring outside of mainland China, one each in Hong Kong SAR and the Philippines. According to the Taiwan Centers for Disease Control, as of 12th February 2020, there were 45167 cases of COVID-19 reported from 28 countries/region and 1115 (2.5%) of patients had died. Among the 45167 cases, most were found in main-land China (n = 44653) and the reported mortality was 2.5% (n = 1113).

**Biochemical mechanism of human Coronaviruses**

All Coronaviruses contain specific genes in ORF1 downstream regions that encode proteins for viral replication, nucleocapsid and spikes formation. The glycoprotein spikes on the outer surface of Coronaviruses are responsible for the attachment and entry of the virus to host cells (Figure 1). The receptor-binding domain (RBD) is loosely attached among virus, therefore, the virus may infect multiple hosts. Other Coronaviruses mostly recognize amino peptidases or carbohydrates as a key receptor for entry to human cells while SARS-CoV and MERS-CoV recognize exopeptidases. The entry mechanism of a Coronavirus depends upon cellular proteases which include, human airway trypsin-like protease (HAT), cathepsins and transmembrane protease serine 2 (TMPRSS2) that split the spike protein and establish further penetration changes. SARS-CoV employs dipeptidyl peptidase 4 (DPP4), while HCoV-NL63 and SARS-CoV-2 require angiotensin-converting enzyme 2 (ACE2) as a key receptor. SARS-CoV-2 possesses the typical Coronavirus structure with spike protein and also expressed other polyproteins, nucleoproteins, and membrane proteins, such as RNA polymerase, 3chymotrypsin-like protease, papain-like protease, helicase, glycoprotein, and accessory proteins. The spike protein of SARS-CoV-2 contains a 3-D structure in the RBD region to maintain the van der Waals forces. The 394 glutamine residues in the RBD region of SARS-CoV-2 is recognized by the critical lysine 31 residue on the human ACE2 receptor. The entire mechanism of pathogenicity of SARS-CoV-2, from attachment to replication is well mentioned in Figure 4.

**Genomic variations in SARS-CoV-2**

The genome of the SARS-CoV-2 has been reported over 80% identical to the previous human Coronavirus (SARS-like bat CoV). The Structural proteins are encoded by the four structural genes, including spike (S), envelope (E), membrane (M) and nucleocapsid (N) genes. The orf1ab is the largest gene in SARS-CoV-2 which encodes the pp1ab protein and 15 nsps. The orf1a gene encodes for pp1a protein which also contains 10 nsps. It is also reported that Spike glycoprotein of the Wuhan Coronavirus is modified via homologous recombination. The spike glycoprotein of SARS-CoV-2 is the mixture of bat SARS-CoV and a not known Beta-CoV. In a fluorescent study, it was confirmed that the SARS-CoV-2 also uses the same ACE2 (angiotensin-converting enzyme 2) cell receptor and mechanism for the entry to host cell which is previously used by the SARS-CoV. The single N501T mutation in SARS-CoV-2’s Spike protein may have significantly enhanced its binding affinity for ACE2.

**Spread, control and prevention of COVID-19**

A person gets infected through close contact with a carrier of the virus and also through coughing and sneezing. Generally, corona virus spreads via air-borne zoonotic droplets. Virus was replicated in ciliated epithelium that caused cellular damage and infection at infection site. According to a study published in 2019, Angiotensin converting enzyme 2 (ACE2), a membrane exo-peptidase in the receptor used by corona virus in entry to human cells.
According to a report published on 24th January 2020\textsuperscript{54}, corona virus infected patients had common features such as fever, cough, and fatigue while diarrhea and dyspnea were found to be as uncommon feature. Many of the patient reported bilateral abnormalities. Corona virus was isolated from bronchoalveolar lavage fluid in China in 2020. It is also detected in blood samples. Till now, corona virus was not confirmed in feaces and urine sample of patient.

To decrease the damage associated with COVID-19, public health awareness and infection control measures are urgently required to limit the global spread of the virus\textsuperscript{59}. Experience from the early phase of SARS-CoV-2 pneumonia strongly highlighted that travel history, rather than chest radiography, is of paramount importance for early detection and isolation of SARS-CoV-2 pneumonia cases. It is essential to limit human-to-human transmission in order to reduce secondary infections among close contacts and health-care workers and to prevent transmission amplification events and further international spread from China. Based on previous experience of management of MERS and SARS infections, the WHO recommend infection control interventions to reduce the general risk of transmission of acute respiratory infections, including avoiding close contact with people suffering from acute respiratory infections, frequent hand-washing especially after direct contact with ill people or their environment, and avoiding unprotected contact with farm or wild animals\textsuperscript{60}. Moreover, people with symptoms of acute respiratory infection should practice cough etiquette, which is to maintain distance, cover coughs and sneezes with disposable tissues or clothing, and wash hands, and within healthcare facilities enhanced standard infection prevention and control practices are recommended in hospitals, especially in emergency departments.

**Therapeutic strategies so far**

2019-nCoV invades through the respiratory mucosa and infects other cells, inducing cytokine storm systemically\textsuperscript{61}. Some patients may progress rapidly with ARDS, disseminated intravascular coagulation (DIC), septic shock, and eventually multiple organ failure\textsuperscript{62}. Therefore, early identification and timely treatment of critical cases is of crucial importance. Evidence-based therapy and supportive care in ICU (Intensive Care Unit) is the mainstay for the management of severe and life-threatening illness of COVID-19. The severe and critical illnesses with COVID-19 should be treated in ICU in the hospital with nosocomial infection control. Strict volume
management, multi-organ function evaluation, critical care of the nutritional assessment/appropriate nutritional support is essential for these patients in ICU. In addition, attention should be paid to bedbound patients to prevent deep vein thrombosis.

**Proposed therapeutic measures**

At present, there is no drug to treat COVID-19 cases. But the under listed are proposed drugs being tried for the treatment of COVID-19.

**Multiple organ functional support and nutrition strategy**

According to the latest epidemiological report, the incidence of the critically ill patients to develop multiple organ dysfunction syndromes is up to 11%. COVID-19 may be combined with other organ injuries, including liver injury, cardiac dysfunction, coagulopathy, which may need the routine functional support for critically ill patients in ICU. Moreover, all the critically ill patients with COVID-19 admitted into ICU have negative nitrogen balance and malnutrition, which has been considered as a contributing factor to the emergence of viral infectious diseases. Therefore, appropriate nutritional strategy is pivotal for the treatment of critical illnesses when necessary.

**Antiviral therapy**

At present, there is no antiviral treatment with confirmed effectiveness for COVID-19. Available drug options that come from the clinical experience of treating SARS, MERS and other previous influenza virus have been used for the treatment of COVID-19 patients. Although these antiviral drugs are promising in the treatment of COVID-19, it should be kept in mind that: (1) the adverse effects of the drugs need to be monitored, (2) the effects of these drugs on critically ill patients still need to be clarified, (3) the potential mutation of the Coronavirus may lead to the drug resistance of the virus.

**Remdesivir**

Remdesivir (GS-5734) is a new nucleoside analog and has been recognized as a potential and promising antiviral drug against a wide array of RNA viruses, including SARS/MERS-CoV. It is currently under clinical development for the treatment of Ebola virus infection. Remdesivir potentially inhibits the RNA-dependent RNA polymerase from MERS-CoV, reduces virus replication, decreases the virus titer in mouse lungs infected with MERS-CoV, and improves the lung tissue damage. The antiviral activity of Remdesivir and IFN-beta was found to be superior to that of the combination of lopinavir/ritonavir and IFN-beta against MERS-CoV. A randomized, controlled trial has reported that the prolonged use of Remdesivir in the treatment of Ebola virus disease (EVD) is safe, and no adverse events have been observed. As a candidate drug that has not been approved, information about the side effects of Remdesivir has not been reported yet. At present, two randomized, controlled, double-blind clinical trials are ongoing to evaluate the efficacy and safety of Remdesivir (200mg loading dose on Day 1, followed by 100mg I.V. once-daily maintenance dose for 9days) in hospitalized patients with mild/moderate or severe COVID-19 respiratory disease. The results of these clinical trials may open the window for effective antiviral therapy for such an epidemic infectious disease.

**Chloroquine and hydroxychloroquine**

It is known that angiotensin-converting enzyme-2 (ACE2) as a membrane protein is a functional receptor of SARS-CoV and it can facilitate virus entry into the cells by binding to the spike (S) protein of the virus, which mediates the fusion of viral and host membranes. Therefore, it may be of importance to block the binding of S protein to ACE2 to treat viral infection, such as SARS-CoV. Chloroquine is a 9-aminoquinoline known since 1934, the sulfate and phosphate salts of which have both been commercialized as widely used antimalarial and autoimmune disease drugs. Chloroquine also shows broad-spectrum antiviral effects. It was found to be a potent inhibitor of SARS-CoV infection due to its inhibitory effects on ACE2. It has been demonstrated that 2019-nCoV enter the epithelial cells of oral mucosa via the essential receptor ACE2, and Chloroquine can function at both entry and post-entry stages of
2019-nCoV infection. Besides the antiviral activity, Chloroquine has an immune-modulating activity, which may synergistically enhance its antiviral effects in vivo. Recently, Wang et al. have demonstrated that Chloroquine is highly effective in the control of 2019-nCoV infection in vitro and is suggested to be assessed in human patients suffering from COVID-19. In addition, the results from more than 100 COVID-19 patients have indicated that Chloroquine phosphate is superior to the control treatment in inhibiting the exacerbation of pneumonia, improving lung imaging, promoting virus negative conversion, and shortening the disease course. However, attention should be paid to the potential detrimental effects of Chloroquine observed in previous attempts to treat viral diseases. At present, the clinical trial to evaluate the efficacy and safety of Chloroquine in the treatment of COVID-19 is ongoing. The use of Chloroquine in the treatment of COVID-19 should refer to the most recent announcements if any. In addition, hydroxychloroquine is a 4-aminoquinoline derivative antimalarial drug. Hydroxychloroquine is an immunosuppressive drug with mature clinical application in the treatment of rheumatic immune diseases such as rheumatoid arthritis and systemic lupus erythematosus. It has been found to be more potent than Chloroquine in inhibiting 2019-nCoV in vitro. Hydroxychloroquine sulfate 400mg given twice daily for 1 day, followed by 200 mg twice daily for another 4 days is recommended in the treatment of COVID-19. At present, the clinical evaluation of hydroxychloroquine in the treatment of COVID-19 is in progress, which might shortly provide preliminary results about the effectiveness of hydroxychloroquine.

**Chinese traditional medicine**

In 2003, Chinese traditional medicine was used to prevent and treat SARS. In 2009, during the pandemic of H1N1 influenza, the Traditional Chinese Medicine of China issued a Chinese traditional medicine prevention program, which included several Chinese herbal medicine formulae for the prevention of infection of adults and children. ShuFengJieDu capsules and Lianhuaqingwen capsules have also played a role in the prevention and treatment of new respiratory infectious diseases such as influenza A (H1N1). Some studies have confirmed that Yupingfeng powder has antiviral, anti-inflammatory and immunoregulatory effects. A multicenter, large-scale, randomized trial found that Yinqiao powder plus another heat-clearing formula could reduce time for fever resolution in patients with the H1N1 influenza virus infection. It is suggested that high-risk populations exposed to COVID-19 patients, including medical staff, family members, and other people who are in close contact with COVID-19 patients, as well as residents living in COVID-19 outbreak areas, might benefit from taking Chinese traditional medicine formulae for prevention. However, the efficacy and safety of these Chinese traditional medicine formulae in COVID-19 need to be further confirmed by clinical trials.

**Methods**

An extensive search spanning from December 2019 to May 2020 was conducted in Medline, EMBASE, Elsevier, Lancet, BioMed Central, PubMed and Cochrane Library databases. The following search terms were used in all possible combinations: (Coronavirus disease 2019” [MeSH] OR “2019 novel coronavirus” [MESH] OR “SARS-CoV-2” [MESH] OR “COVID-19” [MESH] OR “2019-nCov” [MESH], limiting to human subjects. The titles and abstracts of relevant studies were reviewed, and full-text articles were obtained for detailed evaluation which allowed for eligible studies to be included in the systematic review.

**Inclusion and exclusion criteria**

The inclusion criteria were the following: clinical trials, patients with laboratory-identified SARS-CoV-2 infection, treatment outcomes, epidemiological, clinical and laboratory characteristics. The exclusion criteria were absence of clinical characteristics, treatment outcomes, clinical experience and case reports.

**Data extraction and quality assessment**

Two steps were involved in the data extraction process. First, duplicates were removed, and the
remaining articles were screened by their titles and abstracts for relevance. Second, full text articles were reviewed by two of the authors (BOA and EHA) to assess their conformity with the study inclusion criteria. The research domains, publishing dates, journal language, authors’ affiliations, as well as methodological characteristics were analyzed respectively. All findings and statements regarding the outbreak in this review are based on published information as listed in the references.

Conclusion
The outbreak of COVID-19 has become a clinical threat to the general population and healthcare workers worldwide. However, knowledge about this novel virus remains limited. The effective option of antiviral therapy and vaccination are currently under evaluation and development. What we can do now is aggressively implement infection control measures to prevent the spread of SARS-CoV-2 via human-to-human transmission. Public health authorities should keep monitoring the situation, as the more we learn about this novel virus and its associated outbreaks, the better we can respond.

Contributions of Authors
All authors contributed immensely to this review article. Babatunde O. Adetuyi made substantial contributions to conception and design; Peace A. Olajide and Elukunbi H. Awoyelu participated in acquisition of data and drafting of the manuscript; Oluwatosin A. Adetuyi and Oluwaseun A. Adebisi revised the article critically for important intellectual content; and the Julius K. Oloker gave a final approval of the version to be submitted.

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References

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