Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
Nanoparticles as an effective drug delivery system in COVID-19

Neehasri Kumar Chowdhury\textsuperscript{a}, Deepika\textsuperscript{b}, Reshma Choudhury\textsuperscript{c}, Gaurav Ambadas Sonawane\textsuperscript{d}, Shankar Mavinamar\textsuperscript{e}, Xiaoming Lyu\textsuperscript{f}, Ramendra Pati Pandey\textsuperscript{g,}\textsuperscript{h,}, Chung-Ming Chang\textsuperscript{h,}\textsuperscript{*}

\textsuperscript{a} Department of Zoology, Gauhati University, Guwahati, Assam, India
\textsuperscript{b} Department of Biotechnology, Deenbandhu Chhoturam University, and Technology, Sonapat, Haryana, India
\textsuperscript{c} Department of Biotechnology, Royal Global University, Gauhati, Assam, India
\textsuperscript{d} Department of Pharmacy, Sanjivani College of Pharmaceutical Education and Research, Kopargaon, India
\textsuperscript{e} Department of Botany, Karnataka State Akkamahadevi Women's University, Vijayapura, India
\textsuperscript{f} Laboratory Medicine, The Third Affiliated Hospital, Southern Medical University, No.183 West Zhongshan Avenue, Guangzhou, China
\textsuperscript{g} Department of Biotechnology, SRM University, Delhi-NCR, Sonapet, India
\textsuperscript{h} Master & Ph.D. program in Biotechnology Industry, Chang Gang University, No.259, Wenhua 1st Rd., Guishan Dist., Taoyuan City 33302, Taiwan, ROC.

\textsuperscript{*} Corresponding authors.
\textsuperscript{E-mail addresses: neehasri1234@gmail.com (N.K. Chowdhury), deepikagahlot51@gmail.com (Deepika), reshmachoudhury198@gmail.com (R. Choudhury), sonawanegaurav8888@gmail.com (G.A. Sonawane), shankarn050@gmail.com (S. Mavinamar), xiaomlyu@smu.edu.cn (X. Lyu), ramendra.pandey@srmuniversity.ac.in (R.P. Pandey), cmchang@mail.cgu.edu.tw (C.-M. Chang).

https://doi.org/10.1016/j.biopha.2021.112162
Received 3 August 2021; Received in revised form 2 September 2021; Accepted 3 September 2021
Available online 9 September 2021
0753-3322/© 2021 The Authors. Published by Elsevier Masson SAS. This is an open access article under the CC BY-NC-ND license

1. Introduction

A globally spreading disease known as a novel severe acute respiratory syndrome called coronavirus 2 (SARS-CoV-2), abbreviated as COVID-19 pandemic starts spreading from Wuhan in China (as data suggested). On 30th January 2020, WHO declared COVID-19 as a pandemic. The world is dealing with the worst era of the pandemic at a very high infectious and death rate up to 6.89% because of the COVID-19 (data till April 27). The deadliest coronavirus belongs to the Nidovirales order, including Coronaviridae, Arteriviridae, Mesoniviridae, and...
Roniviridae families. Coronaviruses belong to positive-sense mRNA virus and may be characterized by a usually large RNA genome, club-like projections from the surface, and a very unique replication strategy which also shows a mutation in some strains and lethal to human beings [21]. An effective vaccine is still not available to treat this disease hereby it is very important to deliver the drug at a suitable place at right time to reduce the mortality rate and cure infected people [66]. Nanoparticles show a remarkable difference in the culture medium of coronavirus before and during infection with coronavirus. Nanoparticles also show a remarkable inhibition activity or viral life cycle i.e. 55 h [4].

Nanotechnology can be defined as the study, manipulation, control, design, synthesis, and creation of different systems through control of matter at nm (nanometer scale) 1–100 nm where 1 nm = 10^{-9} m i.e. at a molecular and atomic level, utilization of novel phenomenon and properties at that scale [58]. Nanoparticle drug formulations play an important role in pulmonary drug delivery. In dispersed liquid droplets form and as well as in dry powder form different nanoparticles complex shows advantageous result involving the high potential for drug delivery (intracellular), and enhancing dissolution properties makes very un-courageous results to deliver the drug through the lungs [5]. Modern advances of science in drug delivery are very focused upon polymers (chain of monomers) and their types as they exert salient features and distinct biological functions.

The latest approaches in intracellular drug delivery include various polymer-drug conjugates, protein polymer, and stimuli-responsive polymer systems of polymers in drug delivery [39]. Biopolymeric nanoparticles (chitosan and gelatin) can be used to deliver different drugs e.g. gelatin nanoparticles possessing hydrophobicity and hydrophilicity used for delivery of different drugs including doxorubicin [36] and ophthalmic drug [19]. Similarly, to deliver the drug in the case of COVID-19. The loading of the drug on the nanoparticle is defined and based upon the amount of drug per mass of the polymer (usually mg drug per mg polymer or moles of drug per mg polymer). Different techniques can use for analysis and quantification including HPLC, gel filtration, chromatography, spectroscopy, etc.

2. Characteristics of biopolymeric nanoparticles

In the field of Nanotechnology and Nanomedicine, there are large numbers of nanomaterials that are used to deliver drugs having properties like improving solubility, extending formulation’s action, different degrees of lipophilicity, or hydrophilicity, less toxicity. Nanoparticles delivery systems possess different physicochemical properties such as surface properties, shape, size, treatment efficacy, drug system release, and loading, etc. Surface properties of Nanoparticles influenced biocompatibility, biodistribution, and pharmacokinetics of drug molecule [28].

Nanoparticle shape influenced cellular internalization of nanoparticles and biodistribution determination [27]. The size of the nanoparticle can also play a major role in influencing protein adsorption, cellular internalization mechanism, and biodistribution28. Controlled drug release: A significant release of drug cargos including understanding matrix erosion, diffusion, and desorption of linked drug [80]. Drug loading can reduce toxicity from the carrier by reducing the amount of drug carrier used [78]. Biopolymers can be defined as biomolecular monomeric units which are covalently attached to form a large molecule [45]. They are more feasible and suitable as they are not carcinogenic [20]. There are different kinds of biopolymers that can be used in effective drug delivery and preparation of biodegradable nanoparticles including heparin, soluble starch, gelatin, PVP(Polyvinyl-pyrollidone), PVA(Poly-Vinyl Alcohol), cellulose, gelatin, chitosan and so on [45].

Out of all these Chitosan is a much focused and researched biopolymeric nanoparticle. Chitosan is a functional amino polymer having Chitin as a precursor and second most commonly found polysaccharide in nature after cellulose [78]. Chitosan-based NPS have been various applications in different diseases including the brain, ocular infections, pulmonary diseases, gastrointestinal diseases, and cancer [44]. Besides oral, nasal, pulmonary, or vaginal route for drug delivery mucosal route is more feasible and attention gaining part with low toxicity, appropriate physical properties, and good muco-adhesion [54].

Chitosan can deliver drugs by following mechanisms (i) swelling (ii) Diffusion (iii) Erosion. In pulmonary drug delivery, drugs can directly deliver to the lungs which may provide a large surface area which added as an advantage for chitosan nanoparticles hence can be used in the treatment of COVID-19 [56]. The various layer of the lungs including the epithelial lining, mucosal layer, epidermal cells, and branchias mucosal layer can limit drug delivery via lung [57]. Bioavanta Bosti, developed Chitosan polymeric technology in COVID-19 patients develops Novo-chizol [56].

3. Synthesis of biopolymeric nanoparticles

There are various methods used for the synthesis of the bio-nanoparticle-created DDS, for example, spray-drying [34], desolvation [24], electro spraying [6], layer-by-layer self-assembly surfactant Assemblies [64], supercritical fluid extraction [59,74], microemulsion [8, 36] and freeze-drying [32,75]. Among all the listed methods, all method has its advantages and drawbacks also. Between these, desolvation is the simple method for the manufacturing of protein-based nanoparticles by solvents such as ethanol or acetone. In this desolvation method, biopolymeric nanoparticles are manufactured by the mixing of solvents in an aqueous solution by constant stirring. The dimensions of the nanoparticles can be maintained by controlling the flow rate of the amount or volume of a desolvating agent [33,69]. Electrospray technique is a moderate, single step and multipurpose technique with more yield and reliability. To produce particles from size nanometer/micrometer range it uses the electrostatic force to break liquid from a macroscopic mass [42,46]. Freeze-drying is an appropriate method for the compounds which are heat and pressure-sensitive, dried porous particles are obtained when solvents are sublimed. But this method has a drawback, it is a time-consuming and costly method, and the obtained particle having size is generally larger [38,67]. The new layer-by-layer self-assembly method multilayer can be useful for sequential multilayer film formation, which is completed from side to side hydrophobic, electrostatic interactions, and hydrogen bonding are among the films. The nanoscopic features like thickness, surface characteristics, and composition for a film can be achieved by interactions that help to deposit alternate layers of oppositely charged biomaterials and supply precise control to create [30,41]. Finally, by using emulsifiers or surfactants micro-emulsions can be synthesized or prepared by dispersing a biopolymer in two immiscible phases of liquids. Nanomaterials achieved by this method are normally optically transparent, they have high drug solubilizing property, isotropic, and thermodynamically stable (Fig. 1) [11, 36],

4. Role of nanoparticles

Chitosan NP ( nanoparticles) displayed a stamped propensity to aggregate in various tumors. One of the potential explanations behind this marvel might be the brokenness of tumor vasculature. Nano-sized particles can be controlled intravenously because the distance across the littlest blood hair is around 4 µm. Chitosan polymer groups significantly lower harmfulness than poly-L-lysine and PEI is less immuno-genic and needs mutational potential. It upgrades the vehicle of medication across cell film. To build the transfection efficiency, cells focusing on ligands were joined to the chitosan particles. Chitosan-TPP nanoparticles show a lot of potential as reasonable vector contenders for more secure and financially savvy siRNA conveyance. The chitosan nanoparticles have additionally been utilized as non-viral vectors for the quality conveyance or as conveyance transporters for protein particles. Chitosan has broadly been investigated for the application for DNA
mucosal immunizations. A chitosan-based DNA flu antibody has been created by Illum et al. that indicated a high counteracting agent level in mice after intranasal organization. The relationship of immunizations to a portion of the particulate frameworks as nanoparticles has appeared to upgrade the antigen take-up by mucosal lymphoid tissues, in this manner actuating solid foundational and mucosal safe reactions against the antigens. Since chitosan is a low poisonous material, ophthalmic definition dependent on chitosan has displayed a phenomenal resistance after applied chitosan onto the hare’s corneal surface. Other than utilizing chitosan NP to improve tranquilizes transport employing visual, chitosan-covered nanoparticles is likewise used as it displays capacity to upgrade the corneal infiltration [47].

5. COVID-19: history and pathogenicity of coronavirus

From the end of 2019 to yet, the global health sector is passing through a situation that is novel severe acute respiratory syndrome, which is called Coronavirus 2 (SARS-CoV-2) [66]. It is abbreviated as COVID-19 (Coronavirus Disease 2019) [66]. This virus belongs to the family Coronaviridae, the order is Nidovirals and the genus is coronavirus [29] (Fig. 2). These viruses are a group of related RNA viruses and it causes disease in the animals or humans body [29]. This group of viruses causes some respiratory and gastrointestinal infection [50]. It is a zoonotic virus that causes illness and it ranging from the common cold to severe respiratory diseases. Zoonotic virus means these viruses are transmitted from animals to humans (labmanager.com/Lab health and Safety, 2020). Novel coronavirus disease 2019 that is COVID-19, results in a human infection that confirms some signs and symptoms like fever, fatigue, dyspnea, and lymphopenia [50]. In some cases, human infections may lead to some complications like Pneumonia, and cause death [50].

In 1965, Tyrrell and Bynoe found the first cases of coronaviruses in human [29]. It was observed with common cold symptoms, in the human embryonic tracheal organ cultures which are obtained from the respiratory tract of an adult [29]. The first cases of coronavirus (COVID-19) were seen in the Wuhan city of Hubei Province China in December 2019. Before this COVID-19, in 2003, the SARS-CoV virus infected 8098 individuals, and their mortality rate was 9%, across the 26 countries in the World [62]. As of January 30, 2020, there had been 7734 confirmed cases in China, with 90 more cases reported from Taiwan, Thailand, Vietnam, Malaysia, Nepal, Sri Lanka, Cambodia, Japan, Singapore, Republic of Korea, United Arab Emirates, United

Fig. 1. Preparation of nanocomposite using biopolymers.

Fig. 2. Classification of Coronavirus.
States, Philippines, India, Australia, Canada, Finland, France, and Germany. The case fatality rate was calculated to be 2.2% (170/7824) [55] (Fig. 3). It shows that the SARS-CoV-2 (COVID-19) transmission rate is higher than the SARS-CoV because of genetic recombination event at S-protein in the RBD region which is located in SARS-CoV-2 and it may have enhanced for its transmission ability [62].

6. Origin

‘Corona’ name represents the crown-like spikes on the outer surface of the virus so, its name is a coronavirus [62] (Fig. 4). This virus is minute in size and ranges from 65 to 125 nm in diameter. It contains a single-stranded RNA as a nucleic material and its size ranging from 26 to 32kbs in length [62]. The coronavirus family has four subgroups and these are alpha, beta, gamma, and delta coronavirus [62]. The recent COVID-19 virus is a member of the beta group of coronaviruses [62]. Some diseases and their syndrome like, the Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV), H5N1 influenza A, H1N1 2009, and the Middle East Respiratory Syndrome Coronavirus (MERS-CoV) causes acute lung injury and acute respiratory distress syndrome, which cause pulmonary failure and also cause the fatality. These viruses only infect animals until the world occur a severe acute respiratory syndrome (SARS) outbreak, which was caused by SARS-CoV, 2002 in Guangdong, China [83]. After that, another pathogenic coronavirus named the Middle East Respiratory Syndrome coronavirus (MERS-CoV) caused an endemic in the Middle Eastern countries [62]. At the end of 2019, in Wuhan, China; an outbreak of a novel coronavirus emerged that killed and infected various peoples. This virus was named by the Chinese researchers as Wuhan coronavirus or 2019 novel coronavirus (2019-nCoV) [83]. By the international committee on Taxonomy of Viruses (ICTV) named this virus as SARS-CoV-2 and disease named as COVID-19 [17].

7. Pathogenesis

From the end of 2019, COVID-19 has caused mortality and morbidity worldwide [12]. COVID-19 infection is associated with increasing numbers day by day. COVID-19 infectious patients showed higher leucocyte numbers, abnormal respiratory findings, increased levels of plasma pro-inflammatory cytokines [55]. Management of COVID-19 is extremely challenging because some cases showed high infectivity of the virus, but lack of effective antivirals and vaccines, and potentially there exists a large asymptomatic population [12]. According to Genome Sequencing, the COVID-19 (SARS-CoV-2) is partially related with the SARS-CoV (approx. 79% similarity) and MERS-CoV (approx. 50% similarity) [43]. Similar to SARS-CoV, SARS-CoV-2 also uses angiotensin-converting enzyme 2 (ACE2) as its main receptor [43]. It is broadly expressed in vascular endothelium, respiratory epithelium, alveolar monocytes, and macrophages [43]. Here main transmission route is through direct or indirect exposure of the respiratory tract. SARS-CoV-2 shows active replication in the upper respiratory tissues, as demonstrated by the successful live virus which is isolated from throat swabs and detection of viral sgRNA (Sub genomic messenger RNA) in cells of the upper respiratory tract [73].

Some factor contributes for the coagulation disorder in COVID-19 patients [81]. Cytokines like IL-6 could activate the coagulation system and it suppresses the fibrinolytic system [12]. Pulmonary and peripheral endothelial injury causes direct viral attack and it is an equally important inducer of hypercoagulation [81]. With the help of exposure to tissue factors and other pathways, the coagulation system is strongly activated by endothelial cell injury [12]. Moreover, the aggressive immune response is augmented by dysfunctional coagulation [12]. In COVID-19 patients, several levels of anti-cardiolipin and anti-β2GP1 antibodies were detected and needed some additional management strategies [81].

By clinical course, COVID-19 (SARS-CoV-2) infection is divided into three phases i.e.; viremia phase, acute phase (pneumonia phase), and severe or recovery phase [12]. COVID-19 patients with competent immune functions and without some risk factors like old age, co-morbidities, etc., may generate some effective and adequate immune responses for suppressing the virus in the first or second phase without immune overreaction. In contrast to this, COVID-19 patients with immune dysfunction may have a higher risk of failing the initial phase and it becomes a critical type with higher mortality. So, treatment of COVID-19 should be based on the staging of the patients [12]. Management of critical COVID-19 patients are important for reducing the

![Fig. 3. Spreading history of SARS-CoV-2.](image-url)
mortality of the ongoing COVID-19 pandemic. Its key measures lie in monitoring, prevention, and timely intervention. Be prepared for this ongoing pandemic and any other outbreak that might be occurring in the future. It is crucial to understand the pathogenesis of COVID-19 disease. Studies to facilitate the specific therapeutics for control the virus minimize pulmonary injuries or optimize the immune responses [12].

8. Roles of nanoparticles in COVID-19

COVID-19 is an exceptionally deadly respiratory disease that is quickly spreading and has caused universal tension. At present, it is analyzed by identifying the SARS-CoV-2 nucleic corrosive through continuous RT-PCR (qRT-PCR). Nanoparticles are small however it has a huge surface-to-volume proportion, which gives them remarkable and extraordinary highlights in COVID-19. Due to these highlights, nanoparticles have been utilized in the fields of biotechnology, medication, sedate conveyance, sensors, DNA naming and are treated as a scaffold between mass materials. Different delivery systems for drugs are studied under nanotechnology with their possible toxicities. (Table 1).

Point-of-care testing: Point-of-care tests are utilized to get patients without sending tests brought together labs, yielding outcomes without requiring a research facility system to distinguish tainted patients. Horizontal stream antigen location for SARS-CoV-2 is a consideration point, being considered for diagnosing COVID-19. Business sidelong stream tests contain a paper-like film strip covered with two lines: one contains gold nanoparticle-immunizer conjugates; different get the antibodies. At this point, Singular gold nanoparticles are red, and the coupled Plasmon groups cause the arrangement containing the bunch gold nanoparticles to turn blue [71]. Optical biosensor nanotechnology will empower distinguishing the coronavirus in around 30 min legitimately from patients' samples without requiring incorporated research center tests. This innovation could without much of a stretch decide if a patient is contaminated with the coronavirus or the flu infection. This undertaking will possibly be utilized for more than the current pandemic and for treating people. The new biosensor instrument will likewise be utilized to break down different types of coronavirus present in repository creatures.

9. Treatment methodology of COVID-19

We here, sum up the current information to manage potential treatment COVID-19 by Chitosan-associated nanoparticles. It is imperative to alert the peruses that new information refreshing consistently with respect to clinical qualities, analysis, treatment choices, and results for COVID-19. In any case, enhanced steady consideration remains the foundation of treatment and the clinical viability for the resulting specialists is as yet under scrutiny or in clinical trials. Most standing, clinical and preclinical information on antiviral treatment is taken from different Viruses, including SARS-CoV-1 [48], Middle East Respiratory Syndrome, and non-coronaviruses (Ebola) [65]. An affirmed patient of COVID-19 needs total bed rest and steady treatment, guaranteeing satisfactory calorie and water admission to lessen the danger of lack of hydration. Water electrolyte parity and homeostasis need to keep up alongside the checking of indispensable signs and oxygen immersion; keeping respiratory lot unhampered and breathing in oxygen in more extreme cases; estimating blood tally, C-reactive protein, pee test, and other blood biochemical records including liver and kidney work, myocardial catalyst range, and coagulation work as per patient’s conditions. Chest imaging ought to be consistently reevaluated and blood gas investigation ought to be performed when required [1].

Control measures are required for patients with a high fever. Antipyretic medication treatment ought to be acted if the temperature surpasses 38.5 °C. Warm water shower and antipyretic patches are favored as a preventive measure to bring down the temperature. Basic medicaments incorporate ibuprofen orally, 5–10 mg/kg inevitably; acetaminophen orally, 10–15 mg/kg unfaithfully [40].

The odds of hypoxia are expanded as the infection focuses on the lungs. Nasal catheter, cover oxygen ought to be quickly given to the patient. In crisis conditions, Non-intrusive or obstructive mechanical ventilation ought to be given to the patient [61]. Antimicrobial coatings would be a successful measure to control the transmission of COVID-19 at a network level. Metallic nanoparticles are a compelling and mone-tarily reasonable measure to control any pandemic, even COVID-19. Metallic antimicrobial covering plays a multifunctional job and can limit the spread of any lamentable illness in people [53]. A few groups shared their conventions which are summed up in the underneath table (Table 2). Now and again, the gatherings will send reagents or reagent blenders arranged in their research facilities, with or without related

Table 1

| Drug Delivery Systems          | Toxicity Report                      |
|-------------------------------|--------------------------------------|
| Dendrimers                    | Blood toxicity has been found.       |
| Lipid nano systems            | Low toxicity and antigenicity. Presence of cytotoxicity due to surfactant |
| Micelles                      | Low immunogenicity. Presence of cytotoxicity due to surfactant |
| Polymeric nanoparticles       | Required removal of non-degradable polymer. |
| (Chitosan)                    |                                      |
| Engineered nanoparticles      | Presence of Toxicity                 |

Fig. 4. A Structure of Respiratory Syndrome (SARS) Coronavirus.
charges. It is emphatically prescribed to contact the researcher [49].

9.1. Antiviral Drugs

Group of antiviral drugs including interferon α (IFN-α), lopinavir/ritonavir, chloroquine phosphate, ribavirin, and arbidol are therapeutically useful for the Prevention, Diagnosis, and Treatment of Novel Coronavirus-induced Pneumonia by the National Health Commission (NHC) of the People’s Republic of China for tentative treatment of COVID-19 (Table 3).

Favipiravir is a new drug that is under clinical trial for treating COVID-19. On February 15, 2020, China approved it to be a useful drug for treating Novel Influenza. It acts by inhibiting the enzyme RNA-dependent RNA polymerase Apart from being effective for anti-influenza virus, the drug is capable of blocking the replication of flavivirus, alphavirus, bunyavirus, arenavirus, norovirus, and other RNA viruses. Favipiravir is converted into an active phospho-ribofuranosylated form (favipiravir-RTP) in cells and is recognized as a substrate by viral RNA polymerase, thus inhibiting RNA polymerase activity. Therefore, favipiravir may have potential antiviral action on COVID-19, which is an RNA virus [61].

This list was aggregated by the Chinese Pharmaceutical Association, aside from paracetamol, which was included by FIP (Feline Infectious Peritonitis). For the method of reasoning and supporting references for every helpful choice, counsel the first reference (in English), accessible on the FIP committed page. (Chinese Pharmaceutical Association, 2020) Note: This list is for reference just, clinical foundations can make modifications as indicated by their particular conditions (Table 4).

Following are the enlisted antiviral drugs that are possible treatment medicines (Table 5).

The above drugs, have disparate mechanisms of action. Here, a brief insight of all the mechanisms of action of each drug mentioned.

| Country | Institute | Gene targets |
|---------|-----------|--------------|
| China   | China CDC | ORF1ab and N |
| Germany | Charite   | RdRP, E, N   |
| Hong Kong | HKU    | ORF1b-nsP14, N |
| SAR     | National Institute of Infectious Diseases, Department of Virology III | targets, Spike protein |
| Japan   | National Institute of Health | N |
| Thailand | National Institute of Health | N |
| US      | US CDC    | Three targets in N gene |
| France  | Institute Pasteur, Paris | Two targets in RdRP |

9.2. Mechanism of action (remdesivir -GS-5734)

Remdesivir is a monophosphoramidate prodrug of remdesivir-triphosphate (RDV-TP), an adenosine analog that goes about as an inhibitor of RNA-Dependant RNA polymerases (RdRps). Remdesivir-TP contends with adenosine-triphosphate for incorporation into beginning viral RNA chains. When joined into the viral RNA at position I, RDV-TP ends RNA amalgamation (Table 3).

| Type of treatment | Drug name | Dosage form and specifications |
|-------------------|-----------|-------------------------------|
| Antiviral treatment | Recombinant Human interferon α-2a | Injection: 3 million IU, 5 million IU; |
|                   | Interferon α-2b | Injection: 3 million IU, 5 million IU; |
|                   | Lopinavir/ritonavir | Capsule : lopinavir 200 mg, ritonavir 50 mg |
|                   | Ribavirin | Injection: 1 ml: 0.1 g |
| Antimicrobial agents | According to the existing drug list of the medical institution |
| Antipyretic and analgesic treatment | Ibuprofen | Tablet, granules: 0.1 g,0.2 g; Capsule: 0.2 g; Slow release (tablet, capsule): 0.3 g; Suspension: 60 ml:1.2 g, 100 ml:2 g |
| Intestinal microecological preparations | Methylprednisolone | Tablet: 4 mg (Sodium succinate) sterile powder for injection : 40 mg, 500 mg |
| Other gastrointestinal treatment | According to the existing drug list of your medical institution |
| Antitussive treatment | According to the existing drug list of your medical institution |
| Sputum removal treatment | According to the existing drug list of your medical institution |
| Anti-asthmatic treatment | According to the existing drug list of the medical institution |
| Chinese patent medicines | Huoxiangzhengqi | Soft capsule: 0.45 g; Dripping pill: 2.6 g/bag |
|                   | Lianhuaqingwen | Capsule: 0.35 g; Granules: 6 g/bag |
|                   | Shufengjiu | Capsule: 0.52 g |
|                   | Fangfengtongsheng | Concentrated pills: 8 pills equivalent to 6 g herbal slices; Watered pill: 6 g/bag; Granules: 3 g/bag |
|                   | Xuyanping | Injection : 2 ml:50 mg,5 ml:125 mg |
|                   | Xuebijing | Injection : 10 ml |

9.2. Mechanism of action (remdesivir -GS-5734)

Remdesivir is a monophosphoramidate prodrug of remdesivir-triphosphate (RDV-TP), an adenosine analog that goes about as an inhibitor of RNA-Dependant RNA polymerases (RdRps). Remdesivir-TP contends with adenosine-triphosphate for incorporation into beginning viral RNA chains. When joined into the viral RNA at position I, RDV-TP ends RNA amalgamation at position i+3. Since RDV-TP doesn’t cause quick chain termination (i.e., 3 extra nucleotides are consolidated after RDV-TP), the drug seems to proofread by viral exoribonuclease (an...
| Anti-viral Drug     | Structure | Rationale for use                                                                                                                                 |
|-------------------|-----------|--------------------------------------------------------------------------------------------------------------------------------------------------|
| Remdesivir (GS-5734) | ![Remdesivir Structure](image) | Remdesivir is a broad-spectrum antiviral with in vitro activity against coronaviruses [2,9,18,26,35,70,72].                                      |
| Chloroquine       | ![Chloroquine Structure](image) | Chloroquine has in vitro activity against SARS-CoV-2 and has immunomodulating properties [15,70,77].                                               |
| Hydroxychloroquine | ![Hydroxychloroquine Structure](image) | Hydroxychloroquine has in vitro activity against SARS-CoV-2 and may have immunomodulating properties [2,7,15,22,23,60,70].                        |
| Lopinavir         | ![Lopinavir Structure](image) | In vitro and animal model studies show potential activity for other coronaviruses (SARS-CoV and MERS-CoV) [13,14,76].                           |
| Ritonavir         | ![Ritonavir Structure](image) | In vitro and animal model studies show potential activity for other coronaviruses (SARS-CoV and MERS-CoV) [13,14,76].                           |

(continued on next page)
enzyme idea to excise nucleotide analog inhibitors) [2,9,18,26,35,70,72].

9.3. Mechanism of action (chloroquine)

Mechanisms involved inhibition of viral several enzymes or systems such as viral DNA and RNA polymerase, glycosylation of viral protein, virus assembly, new virus particle transport, and virus discharge. Other mechanisms may also involve ACE2 cellular receptor inhibition, acidification at the surface of the cell membrane repressing fusion of the virus, and immunomodulation of cytokine release [2,7,16,22,23,60,70].

9.4. Mechanism of action (hydroxychloroquine)

Mechanisms involved inhibition of viral several enzymes or systems such as viral DNA and RNA polymerase, glycosylation of viral protein, virus assembly, new virus particle transport, and virus discharge. Other mechanisms may also involve ACE2 cellular receptor inhibition, acidification at the surface of the cell membrane repressing fusion of the virus, and immunomodulation [7,16,22,23,60,70].

9.5. Mechanism of action (lopinavir & ritonavir)

Lopinavir and ritonavir may bind to Mpro, a key enzyme of coronavirus replication. This inhibits coronavirus activity [41].

9.6. Mechanism of action (favipiravir)

Favipiravir is an RNA-dependent RNA polymerase (RdRp) inhibitor that inhibits viral transcription [15,63,72].

10. Adverse outcome pathway

The world is facing a continuous struggle with a much larger potential to spread the deadly and highly infectious SARS-CoV-2. The reuse of some prospective antiviral medications is the main area of investigation from a therapeutic point of view. For suitable safety and pharmacokinetic profile in people, some additional medicines with a considerably higher in vitro and in vivo efficacy against COVID-19 were studied. In its creation and use, however, there are other problems. The coronaviruses are variable RNA viruses. New coronavirus varieties with new structures thus appear easily. For these new coronavirus types, conventional medicines may not be as effective or have just weak effects. Further study may be carried out in order to accomplish multimodal therapies for various antiviral medicines concurrently on-site. The creation of these new forms of antiviral medicine distribution is usually more difficult than the standard system for the delivery of medications. The challenges of large-scale production are amongst the few impediments to the development of this promising technology. The optimized manufacture of the nanocarrier system to encapsulate therapy potential utilizing the quality-by-design method in order to highlight the crucial processing factor and content material characteristics is important. Furthermore, the design of the innovative drug delivery is critical to an integrated understanding of COVID-19’s structural morphology, virulence and transmission.

11. Discussion

In global health, the SARS-CoV-2 novel – known as the COVID-19 pandemic – has the most dangerous effect that affects people in every region and community in the world. The development of COVID-19 diagnostics, treatment, prevention and developing vaccine may play a crucial role in nanotechnology based medicines. Moreover it is important to have time for the encounter of infectious diseases and nano-based materials in comparison with other therapies, as they are not linked to long and strict regulatory issues commonly associated with vaccines, to come to the public earlier. Even in small concentrations and shortening response periods to see if an individual is infected, it is necessary to develop a broad antiviral spectrum of disinfectants with acceptable effectiveness. In addition, it is crucial that monotherapy problems are overcome and that the drug’s dose is reduced and its negative effects decreased.

Moreover a solution that can reduce the risk of diseases where the viral load is heavily reduced is the development of surface coating with a relatively small surface energy (which can repel, kill or be anti-adhesion to the pathogens). Nano-based materials have the potential to gain a foothold in the face of the current global public health hazard by encouraging the wide-ranging, integrated approaches that are required to manage and control the COVID-19 outbreak at both the local and international levels.

Chitosan nanoparticles will be the most emergent field to study drug delivery and novel drug discovery. In pulmonary drug delivery, drugs are administered directly to the lungs, offering a wide surface area, which is a benefit for chitosan nanoparticles, allowing them to be utilised to treat COVID-19. Our current studies reveal that nanoparticles may prove very helpful in the delivery of drugs for covid-19 treatment.

12. Conclusion

In recent studies, it has been seen that nanoparticles can prove very helpful in the delivery of drugs for COVID-19. In many cases, it has been observed that patients, where drugs are delivered with the help of nanoparticles, produced very few side effects. Several Drugs have been used to prevent COVID-19 infection including interferon α (IFN-α), lopinavir/ ritonavir, chloroquine phosphate, ribavirin, and arbidol. A new drug Favipiravir was also launched to treat COVID-19 in February 2021. These all drugs should be delivered into the human body at a specific range. Biopolymeric nanoparticles can be used to deliver these kinds of drugs by minimizing the risk of toxicity, with low solubility, and increasing feasible delivery of a drug. Chitosan nanoparticles will be the most emergent field to study drug delivery and novel drug discovery. In pulmonary drug delivery, drugs are administered directly to the lungs, offering a wide surface area, which is a benefit for chitosan nanoparticles, allowing them to be utilised to treat COVID-19. Our current
studies reveal that nanoparticles may prove very helpful in the delivery of drugs for COVID-19 treatment.

13. Future prospects

Nanomedicine science is one of the most exciting fields of research at the present time. The implementation of nanomedicine and of the nano-drug delivery system will certainly remain the future arena of research and development for many years, with the use of a variety of nano-particles in the precise supply of drugs to the affected cells without disturbing the physiology of normal cells. Although the prospects for nanomedicine and the nano-drug system are overwhelmingly clear, their real effect is still very limited on the healthcare system, even in continuing to be demanded for their biodegradability, biocompatibility, carcinogenic /diagnostics. The field is characterized by a new area of drug delivery system will certainly remain the future arena of research science, which only has real research on the subject over two decades and many key fundamental attributes are still unknown. One main area of research for the future is the fundamental marker of diseases with key biological markers which enable absolute focusing without changing the normal cellular process. New natural biomaterials are continuing to be demanded for their biodegradability, biocompatibility, readiness for availability, renewability, and low toxicity quality. Nanobiosensors can be used extensively for rapid detection of the viral disease.

CRediT authorship contribution statement

NK, D, RC, GAS, SM, RPP, and CMC were involved in the study’s conceptualization and design. The material was organized and the initial draft of the manuscript was written by NK, D, RC, GAS, SM, RPP, and CMC. NKC, D, RC, GAS, SM, RPP, and CMC contributed to the manuscript by writing portions and assisting with manuscript editing and formatting. All authors helped to revise the manuscript, read it, and approved the final version.

Conflict of interest statement

The author declares no conflict of interest, financial or otherwise.

Acknowledgements

This work was supported through the industry-academia collaboration project, VTR Inc CGU, R.O.C., project grant (SCRPD110221).

References

[1] E. Abbasi-Oshaghi, F. Mirzaei, F. Farahani, I. Khodadadi, Biaryzahnia, Diagnosis and treatment of coronavirus disease 2019 (COVID-19): laboratory, PPR, and chest CT imaging findings, Int J. Surg. 79 (2020) 143–153, https://doi.org/10.1016/j.ijsu.2020.05.018. PMID: 32422384; PMCID: PMC7227548.
[2] M.L. Agostini, E.L. Andrews, A.C. Sims, P.G. Graham, T.P. Sheehan, X. Lu, R. Jordan, Coronavirus susceptibility to the antiviral desiravir (GS-5734) is mediated by the viral polymerase and the proofreading exoribonuclease, mBio 9 (2018).
[3] P. Bablano, Y. Shamsi, P. Kapoor, M. Sharma, Y. Arora, Inflamed site-specific drug delivery system based on the interaction of human serum albumin nanoparticles with myeloperoxidase in a murine model of experimental colitis, Eur. J. Pharm. Biopharm. 125 (2018) 141–182.
[4] W.-Z. Jiang, Y. Cai, H.-Y. Li, Fish-based spray-dried mucocarethosis microspheres for sustained oral mucosal drug delivery, Powder Technol. 312 (2017) 124–132.
[5] W.-C. Ko, M.-H. Rolen, N.-Y. Lee, P.-L. Chen, C.-T. Huang, P.-I. Lee, R.H. Hsieh, In vitro susceptibility of 10 respiratory syncytial virus strains to ribavirin, J. Genet. Genom. Yi Chuan Xue Bao 47 (2) (2020) 119–121.
[6] W. Liang, A.Y. Ayhan, M.Y. Chow, F.F. Lo, Y. Qiu, P.C. Kwock, J.K. Lam, Spray freeze drying of small nucleic acids as inhaled powder for pulmonary delivery, Asian J. Pharm. Sci. 13 (2) (2018) 165–172.
[7] W.-B. Liang, B. Kryscio, D.R. Kriciuk, B.V. Slaughter, N.A. Peppas, Polymers for drug delivery systems, Annu. Rev. Chem. Biomol. Eng. 1 (2010) 149–173.
[8] S. Kupper, I. Klosowska-Chomiczewska, P. Szumala, Collagen and hyaluronic acid hydrogel in water-in-oil microemulsion delivery systems, Carbohydr. Polym. 175 (2017) 347–354.
[9] S. Luo, T. Okubo, M. Nangrejo, M. Edirisinghe, Preparation of polymeric nanoparticles by novel electrospray nanoprecipitation, Polym. Int. 64 (2) (2015) 187–192.
[10] S. Mehta, D.F. McAuley, M. Brown, K. Shuke, P. Sanchez, J.J. Manson, H.A. Mehta, Potential inhibitors against 2019-nCoV coronavirus M protein from clinically approved medicines, J. Genet. Genom. Yi Chuan Xue Bao 47 (2) (2020) 119–121.
[11] S.P. Callender, J.A. Mathews, K. Kobernyk, S.D. Wettig, Microemulsion utility in polymerase, Antivir. Res. 169 (2019), 104541.
M. Wang, R. Cao, L. Zhang, X. Yang, J. Liu, M. Xu, G. Xiao, Remdesivir and B. Von Storp, A. Engel, A. Boeker, M. Ploeger, K. Langer, Albumin nanoparticles F. Ungaro, O. Catanzano, I. d K. Shiraki, T. Daikoku, Favipiravir, an anti-influenza drug against life-threatening A. Savarino, L. Di Trani, I. Donatelli, R. Cauda, A. Cassone, New insights into the E. Rytting, J. Nguyen, X. Wang, T. Kissel, Biodegradable polymeric nanocarriers for C.A. Ruge, J. Kirch, C.-M. Lehr, Pulmonary drug delivery: from generating aerosols A. Rampino, M. Borgogna, P. Blasi, B. Bellich, A. Cesaro, New insights into the challenges, Nanomaterials 10 (5) (2020) 852. P.K. Rai, Z. Usmani, V.K. Thakur, V.K. Gupta, Y.K. Mishra, Tackling COVID-19 infection, 26 May 2020., World J. Pedia (2019). N.K. Chowdhury et al. monomers, Nature 531 (7594) (2016) 381–230. F. Quaglia, Microparticle-embedded fibroin/alginate beads for prolonged local challenges, Nanomaterials 10 (5) (2020) 852. I.S. Chronakis, Development of electrosprayed mucoadhesive chitosan nanoparticles, Carbohydr. Polym. 190 (2018) 240–247. K. Naggal, S.K. Singh, D.N. Mishra, Chitosan nanoparticles: a promising system in novel drug delivery, Chem. Pharm. Bull. 59 (11) (2011) 1423–1430. B.R. O Keefe, B. Giammaretto, D.L. Barnaard, S.R. Shenoy, P.K. Chan, J.B. McMahon, C.L. Wohlford-Lenane, Broad-spectrum in vitro activity and in vivo efficacy of the antiviral protein griffithsin against emerging viruses of the family Coronaviridae, J. Virol. 84 (5) (2010) 2511–2521. Organization, W.H. (2020). Population-based age-stratified seroepidemiological investigation protocol for coronavirus (COVID-19) infection, 26 May 2020. Retrieved from. R.A. Petros, J.M. DeSimone, Strategies in the design of nanoparticles for therapeutic applications, Nat. Rev. Drug Discov. 9 (8) (2010) 615–627. P.K. Rai, Z. Usmani, V.K. Thakur, V.K. Gupta, Y.K. Mishra, Tackling COVID-19 pandemic through nanocoatings: confront and exactitude, Curr. Res. Green. Sustain. Chem. 3 (2020), 100011. A. Rampino, M. Borgogna, P. Blasi, B. Bellich, A. Cesaro, Chitosan nanoparticles: preparation, size evolution and stability, Int. J. Pharm. 455 (1–2) (2013) 219–228. H.A. Rothen, S.N. Byrzeddy, The epidemiology and pathogenesis of coronavirus disease (COVID-19) outbreak, J. Autoimmun. 109 (2020), 102433. C.A. Ruge, J. Kirch, C.-M. Lehr, Pulmonary drug delivery: from generating aerosols to overcoming biological barriers—therapeutic possibilities and technological challenges, Lancer Respir. Med. 1 (5) (2013) 402–413. E. Rytting, J. Nguyen, X. Wang, T. Kissel, Biodegradable polymeric nanocarriers for pulmonary drug delivery, Expert Opin. Drug Deliv. 5 (6) (2008) 629–639. F. Salamandra-Buentello, D.L. Persad, D.K. Martin, A.S. Daar, P.A. Singer, Nanotechnology and the developing world, PLoS Med 2 (5) (2005) 97. A. Salerno, L. Verdolotti, M. Rauci, J. Saurina, C. Domingo, R. Lamanna, M. Lavorgna, Hybrid gelatin-based porous materials with a tunable multiscale morphology for tissue engineering and drug delivery, Eur. Polym. J. 99 (2018) 230–239. A. Savarino, L. Di Trani, I. Donatelli, R. Cauda, A. Cassone, New insights into the antiviral effects of chloroquine, Lancet Infect. Dis. 6 (2) (2006) 67–69. K. Shen, Y. Yang, T. Wang, Diagnosis, treatment, and prevention of 2019 novel coronavirus infection in children: experts’ consensus statement [published online February 7, 2020], World J. Pediatr. (2019). M.A. Sheereen, S. Khan, A. Kazmi, N. Bashir, R. Siddique, COVID-19 infection: origin, transmission, and characteristics of human coronavirus, J. Adv. Res. 24 (2020) 91–98. K. Shiraki, T. Daikoku, Favipiravir, an anti-influenza drug against life-threatening RNA virus infections, Pharmacol. Ther. 209 (2020), 107512. Y. Shon, H. Kim, H.S. Hwang, E.S. Bae, T. Eom, E.J. Park, B.S. Shim, A nanostructured cell-free photosynthetic biocomposite via molecularly controlled layer-by-layer assembly, Sens. Actuators B: Chem. 244 (2017) 1–10. D. Siegel, H.C. Hui, E. Dorrerfield, M.O. Clarke, K. Chum, L. Zhang, B. Ross, Discovery and Synthesis of a Peptide Library: Predatory Proteinase Inhibitors for Detection of 2,1-(triazin-4-aminos) Adenine C-Nucleoside (GS-5734) for the Treatment of Ebola and Emerging Viruses, ACS Publications, 2017. V.S. Sivasankarappillai, A.M. Pillai, A. Rahdar, A.P. Sobha, S.S. Das, A. Citropolous, G.Z. Kyzas, On facing the SARS-CoV-2 (COVID-19) with combination of nanomaterials and medicine: possible strategies and first challenges, Nanomaterials 10 (5) (2020) 852. F. Ungaro, O. Catanzano, I. d Angelo, L. Diaz-Gomez, A. Concheiro, A. Miro, F. Quaglia, Microparticle-embedded fibrous/alginic beads for prolonged local release of simvastatin hydroxyacid to mesenchymal stem cells, Carbohydr. Polym. 175 (2017) 645–653. B. Von Storp, A. Engel, A. Boeker, M. Ploeger, K. Langer, Albumin nanoparticles with predictable size by desolvation procedure, J. Microencapsul. 29 (2) (2012) 138–146. M. Wang, R. Cao, L. Zhang, X. Yang, J. Liu, M. Xu, G. Xiao, Remdesivir and chloroquine effectively inhibit the recently emerged novel coronavirus (2019- nCoV) in vitro, Cell Res. 30 (3) (2020) 269–271. A. Waris, M. Ali, A.U. Khan, A. Ali, A. Baset, Role of nanotechnology in diagnosing and treating COVID-19 during the Pandemic. T.K. Warren, R. Jordan, M.K. Lo, A.S. Ray, R.L. Mackman, V. Soloveva, H.C. Hui, Therapeutic efficacy of the small molecule GS-5734 against Ebola virus in rhesus monkeys, Nature 531 (7594) (2016) 381–385.