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.

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Respiratory viruses represent a severe public health risk worldwide, and the research contribution to tackle the current pandemic caused by the SARS-CoV-2 is one of the main targets among the scientific community. In this regard, experts from different fields have gathered to confront this catastrophic pandemic. This review illustrates how nanotechnology intervention could be valuable in solving this difficult situation, and the state of the art of Zn-based nanostructures are discussed in detail. For virus detection, learning from the experience of other respiratory viruses such as influenza, the potential use of Zn nanostructures as suitable sensing platforms to recognize the S1 spike protein in SARS-CoV-2 are shown. Furthermore, a discussion about the antiviral mechanisms reported for ZnO nanostructures is included, which can help develop surface disinfectants and protective coatings. At the same time, the properties of Zn-based materials as supplements for reducing viral activity and the recovery of infected patients are illustrated. Within the scope of noble adjuvants to improve the immune response, the ZnO NPs properties as immunomodulators are explained, and potential prototypes of nanoengineered particles with metallic cations (like Zn\(^{2+}\)) are suggested. Therefore, using Zn-associated nanomaterials from detection to disinfection, supplementation, and immunomodulation opens a wide area of opportunities to combat these emerging respiratory viruses. Finally, the attractive properties of these nanomaterials can be extrapolated to new clinical challenges.

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1. Introduction: COVID-19 and its emergence

According to the World Health Organization, the current coronavirus outbreak (COVID-19) has spread worldwide, giving rise to a pandemic with roughly 5 million deaths reported so far. Unfortunately, the pandemic continues to expand and remains without proper treatment. In addition, the accelerated surge of new viral strains such as the so-called Delta variant has led to new records in hospitalization and deceased cases [1]. The etiological agent of COVID-19 is the Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2). This is an enveloped virus, which means that the second layer of glycoproteins surrounds the capsid that contains its single-stranded RNA genome. Its structure comprises the Spike protein (S) (S1 and S2 domains), membrane protein M, envelop protein E, and the nucleocapsid protein N. The viral recognition occurs through the interaction between the host cell receptor Angiotensin-Converting Enzyme 2 (ACE2) and the S protein, then followed by endocytosis and subsequent viral replication (Fig. 1 (a-b)) [2].

Apart from the novel SARS-CoV-2, there are other viruses responsible for causing respiratory diseases such as SARS-CoV, MERS coronavirus (MERS-CoV), syncytial virus, rhinovirus, and influenza virus, whose contagion capacity has caused former pandemic outbreaks (e.g., SARS-CoV, 2003 [3], H1N1, 2009 [4] and MERS, 2012 [5]). Compared with the SARS-CoV-2, the SARS-CoV possess 79.5% of identity and MERS 40%, both being part of the same β-coronavirus class [6]. On the other hand, the influenza virus resembles coronaviruses in its spherical morphology, nanometric size (80–120 nm), and rapid mutagenic capacity. The influenza viruses can be divided into four genera A, B, C, and D, where the principal human infection occurs due to the A and B subtypes. It is also an enveloped virus surrounded by hemagglutinin and neuraminidase glycoproteins, being the first one responsible for host cell recognition [7]. In the case of Human rhinovirus (HRVs), its size is smaller (~30 nm) and, unlike the coronavirus, it has a non-enveloped structure and icosahedral morphology (Fig. 1(c)). HRVs cause most acute respiratory infections worldwide. Thus, understanding how these viruses enter the human cell and their replication mechanisms allow us to extrapolate previous strategies to fight back the ongoing pandemic.

The different mutations that the respiratory viruses undergo can lead to continuous seasonal outbreaks where prophylaxis and early detections play a determining role. The above-mentioned leads to the ongoing necessity of developing novel materials that help us mitigate current and future pandemics. In this regard, the nanomaterials have been used to deal with the SARS-CoV-2 and other viruses, among which we can highlight drug delivery [8], immunomodulatory responses [9], disinfecting [10,11], and biosensing applications [12-22]. For biosensing, several platforms are based on Au, Se, Fe, Graphene, and Zn (for further details, see Table 1 in Section 2). However, one of the elements that have been studied for biosensor platforms is zinc (Zn) due to its excellent optical and electronic properties, with the additional advantage of being cheap and abundant in nature. A variety of Zn-based nanostructures (e.g., ZnO, CdZnSeS/ZnSeS, CdSe/CdS/ZnS, C-ZnO) continue to be extensively investigated for detecting respiratory viruses. For example, in work developed by Thervasahayam et al. [23], ZnO NRs have been functionalized with 1-ethyl-3-(3-dimethyl aminopropyl) (EDC) (linker) and the ACE2 protein to form the Zn-ZnO-EDC-ACE2 species that allows rapid identification and quantification of the virus in 18–60 s [24]. ZnO nanostructures have also been used for the selective detection of other emerging viruses, for instance, H1N1, H5N1, and H7N9 influenza viruses (Section 2b, Table 2); as can be observed, the detection limit for Zn-based nanomaterials is comparable with the reported materials up to date.

On the other hand, one of the principal concerns about the SARS-CoV-2 is its high contagiousness, increasing the mortality rate worldwide. The main transmission, in this case, is through expelled droplets from infected individuals or indirectly by contact with contaminated surfaces. For this reason, Zn/ZnO nanoparticles (NPs) are also used as disinfectants and have been incorporated in commercial products, e.g., in food packing, owing to their low cytotoxicity and high antimicrobial activity [25-27]. Furthermore, the use of nano Zn as disinfectant agents could restrict the SARS-CoV-2 transmission by developing self-sterilized coatings since the virus can remain active over plastic and stainless steel surfaces [28].

Further, COVID-19 symptomatology varies from mild (fever, cough) to severe (pneumonia or acute respiratory distress syndrome (ARDS)) to fatality. Casualties are most common in patients with comorbidities, like obesity [29,30], cardiovascular disease [31], diabetes mellitus [32,33], hypertension [34], and cancer [35] due to inflammatory and immune response complications. In this sense, recent medical research has highlighted Zn as an essential element in this viral infection; e.g., the diseases mentioned above (diabetes and hypertension) are associated with a Zn deficiency [36,37]. Likewise, anemia and dysfunctional taste are also related to low levels of Zn [38,39]. Additionally, it has been found that Zn can improve innate and adaptive immunity in the course of a viral infection. Furthermore, Zn supplementation (based on different nano Zn compounds) can boost the recovery of SARS-CoV-2 patients [40-44] (Fig. 1 (d)). Notably, Zn is involved in multiple human body responses, including the immune system [45], and this explains why it has been employed as a treatment for other pathologies such as pneumonia and diarrhea by rhinovirus [46,47], stunted growth in children [48,49], Wilsons disease [50], and acrodermatitis enteropathica [51] to mention a few.

As in the case of bacteria, viruses are also susceptible to developing resistance to specific antiviral treatments. In this sense, given the remarkable properties of Zn, Zn-based nanomaterials have also been tested against several respiratory viruses, thus demonstrating their potential application as a treatment. The commonly proposed antiviral mechanism suggest the delivery of Zn^{2+}...
ions, which are active against the SARS-CoV-2 due to its capacity to reduce the inflammatory responses by (i) inhibition of NF-κB and modulation of T-cell [52], (ii) decreasing the activity of ACE2, (iii) avoiding the entry into the host cell, and (iv) regulating the interferon α production (INFα) [53]. For instance, a combination of ZnO NPs-aspartic acid reduces the duration of the common flu symptoms caused by rhinovirus and increases the number of CD4+ T individuals infected with the Human Immunodeficiency Virus (patent application number PCT/US01/20579C [54]). The former immunomodulatory properties also suggest using Zn-based nanomaterials as adjuvants to enhance the release of inflammatory cytokines, activating the immune cell.

Different databases such as Web of Science, PubMed, and Journal of the American Medical Association (JAMA) have been used to compile information. To identify the relevant literature, multiple keywords and term combinations were used, including “SARS-CoV-2”, “COVID-19”, “Zn,” “studies” “biosensor,” “treatment,” “completed studies,” “respiratory virus,” “supplement,” and “nano” keywords with filters. The results show an increase in the extent of nanotechnology research compared with 2018, where almost no results were found for vaccination. As shown in Fig. 2 (a), there is an uprising trend in the number of publications linked to the use of nano Zn in supplementation applications or directly associated with the treatment of viruses, thus reflecting the relevance of this material in health-related applications. However, only a tiny fraction of the publications mentioned focus on using this material in a nanostructured form, which demands further research and development in this field. Fig. 2(b–d) shows tree diagrams of the research fields obtained using the Web of Science analysis tool for various keyword combinations. These graphs show the multidisciplinary nature of the bionanotechnological applications and visualize the necessity of an interdisciplinary approach. Finally, the interrelation between the fundamental concepts of the systematic search is outlined in Fig. 2(e), employing a Venn diagram. It is possible to observe that the intersection between nano Zn and COVID-19 is small and yet to be explored in detail.

The present review highlights the state-of-the-art nanostructured Zn-based materials as biosensing platforms, antimicrobial coatings, supplements in the diet, and potential adjuvants in immunomodulation, which are key points to handle the current pandemic and future events related to respiratory virus infections (Fig. 1(f)). We aim to encourage the implementation of nano Zn materials into future treatments and the development of...
| Nanostructure | Interaction mechanism | Response signal | Analyte | Bioselective layer | Synthesis technique | Limit of detection/Linear response range | Reference |
|---------------|-----------------------|-----------------|---------|--------------------|---------------------|------------------------------------------|-----------|
| Au NIs        | DNA hybridization      | PPT* & LSPR*    | RdRp-COVID, ORF1ab-COVID and E genes from SARS-CoV-2 | Hybridization probe (ssDNA) | Self-assembly thermal dewetted of Au thin film | 0.22 pM/ 5–1 pM | [67] |
| Au NPs        | DNA hybridization      | Optical absorbance | SARS-CoV-2 nucleocapsid phosphoprotein (N-gene). | Oligonucleotide probe (ssDNA) | Modified citrate reduction | 0.18 ng mL⁻¹ / 0.2–3.0 ng mL⁻¹ | [12] |
| AuNPs-graphene (Paper-based platform) | DNA hybridization | Electrochemical | SARS-CoV-2 nucleocapsid phosphoprotein (N-gene). | Oligonucleotide probe (ssDNA) | Modified citrate reduction/ Spin coating | 6.9 copies µL⁻¹ | [63] |
| FTO electrode-AuNPs | Antigen-antibody interaction | Electrochemical: Amperometric | Covid-19 spike antigen | Covid-19 monoclonal antibody | Modified citrate reduction | 10 fM / 1 fM–1 µM | [271] |
| Graphene sheet | Protein-antibody interaction | Field-effect transistor | SARS-CoV-2 spike protein | Antibody against SARS-CoV-2 spike protein | Wet-transfer method | Culture: 1.6 × 10¹ pfu mL⁻¹ / clinicsamples: 2.42 × 10² copies mL⁻¹ | [70] |
| rGO nanolakes | DNA hybridization | Electrochemical: Impedance | RDB SARS-CoV-2 spike protein | Antibody against SARS-CoV-2 spike S1 protein | 3D nanolithography / drop-casting process | 16.9 × 10⁻¹⁻¹ M | [71] |
| CdSe QDs      | DNA hybridization      | Fluorescence     | SARS-CoV-2 nucleocapsid phosphoprotein (N-gene). | RNA aptamer | Commercially available CdSe QDs. Invitrogen Corp (QD605). | 0.1 pgmL⁻¹ | [75] |
| Fe₃O₄ MNPs*   | DNA hybridization      | Transcription    | SARS-CoV-2 viral sequence | Carboxylizedpoly (amino ester) | Co-precipitation | 10 copies / 10⁻¹⁻¹ copies | [21] |
| Iron oxide NPs | DNA hybridization      | RT-PCR & magnetic response | C2CA-optimagnetic detection | RDB SARS-CoV-2 sequence | Biotinylated detection probe | Commercially available MNPs. Micromod PartikelterchnologieGmbH. | 0.4 fM | [79] |
| Chemiresistance | Antigen-antibody interaction | Electrochemical: capacitance change | SARS-CoV-2 spike protein | Zinc containing metalloenzyme receptor | Hydrothermal | Patent application number IN2020-41034067 | [23] |

Au NIs: Gold Nanoislands; Au NPs: Gold nanoparticles; FTO: Fluorine doped thin oxide; rGO: reduced graphene oxide; Lanthanide-doped polystyrene nanoparticles; MNPs: Magnetic Nanoparticles; C2CA: Circle-to-circle amplification; QDs: Quantum dots; VOCs: volatile organic compounds; PPT: Plasmonic Photothermal Effect; RBD: Receptor Binding Domain; LSPR: Localized Surface Plasmon Resonance; SPR: Surface Plasmon Resonance; ssDNA: single-strand DNA.

*MNPs application to enhance PCR amplification efficiency.
point-of-care devices, which also demands further research. The results found in the present work point them out as promising candidates to be employed in health emergencies such as emerging respiratory viruses.

2. A summary of nanomaterials for the detection of COVID-19

As stated by many authors [10,15,55-57], one of the main axes of the transversal efforts to confront this global health emergency lies in the efficient, rapid, economic, and scalable development of biosensing platforms for the precise and selective detection of the SARS-CoV-2. Therefore, the fast and accurate detection of infected individuals in the early stages of the disease is vital as it allows prompt treatment and prevents further dissemination.

Currently, the most used techniques for the detection of human coronaviruses include nucleic acid amplification (e.g., reverse transcriptase-polymerase chain reaction (RT-PCR)) and sequencing, lung visualization through computer tomography, chest radiography or ultrasound, cultures, and antibody-antigen immunoassays tests (serological assays) [58]. However, as the magnitude of the pandemic continues to rise, these methods have not been able to cope due to limitations such as time-consuming, expensive instruments required, elaborate sample preparation, and false-positive results [59,60].

Consequently, incorporating specific nanomaterials into emerging technologies such as point-of-care tests and optical, electrochemical, or magnetic biosensors can contribute to the efficient surveillance of SARS-CoV-2.

a. An update on different nanomaterials employed as nanobiosensors

Viruses are a typical example of recurrent nanostructured materials in nature; for instance, in the case of the SARS-CoV-2, it is commonly described as a spherical-like particle with a diameter from 120 to 160 nm [61]. Consequently, the detection of such small biomarkers has been enhanced by implementing nanostructured transducing materials into multiple biosensing platforms. The improvement is mainly attributed to the high surface-to-volume ratio of the nanomaterials that increases the interaction with the biological analyte and results in selective and sensitive detection.

Real-time fluorescence quantitative RT-PCR plus viral gene sequencing are the gold standards for diagnosing SARS-CoV-2 infection [62]. However, the need for quick detection methods is one of the most critical control strategies. Currently, different works have been published in the field of biosensors to detect the new COVID-19.

Among the various nanomaterials used to detect human viruses, Au nanostructures, e.g., NPs and nanorods (NRs), have been used as bio-transducing systems to detect COVID-19 employing different types of bioreceptors [12,13,18,63,64], thereby allowing an accelerated transition of these systems to COVID-19 diagnostic devices (Fig. 3(a)). Wen et al. [65] reports the implementation of Au NPs coated with IgG antibodies against SARS-CoV-2 into a lateral flow immunoassay strip (LFAs) with reasonable specificity and sensitivity. The system allows to detect immune response, customarily manifested two weeks after contracting the infection [66]. On the other hand, a hybridization probe detection approach is reported by Qui et al. to detect the SARS-CoV-2 sequence [67]. A thiol bond attached the bioreceptor to the surface of the Au nanoflakes. Then, the Localized Surface Plasmon Resonance (LSPR) Plasmonic Photothermal (PPT) responses were used to detect the targeted viral sequences with a LOD of 0.22 pM. Discrimination tests performed in multigene mixtures (SARS-CoV/SARS-CoV-2) showed good selectivity towards the SARS-CoV-2 virus due to the localized thermal enhancement of the signal. Other reported detection systems include optical biosensors [18] (Fig. 3(b)), SERS (Surface Enhanced Raman Spectroscopy) platforms [60] (Fig. 3(c)), and Colorimetric [12] sensors, allowing the fabrication of detection kits that enable the user to know the response with the naked eye. Therefore, the development of this kind of Point of Care (PoC) test could offer a viable solution to detect the virus where no infrastructure for molecular tests is available, for example, in distant or marginalized communities.

On the other hand, various carbon nanostructures have proved to be effective biocompatible materials for detecting human viruses [68,69] and specifically the SARS-CoV-2 virus [68,70,71]. A microfluidic electrochemical biosensor for detecting SARS-CoV-2 spike S1 protein employing reduced-graphene-oxide (rGO) nanoflakes as electrode covering material is reported by Azahar et al. [71]. They reported a LOD of $16.9 \times 10^{-15}$ M, good reusability, and no cross-reaction with other antibodies or interfering proteins (Fig. 4(a-e)). Similarly, the novel coronavirus S protein was targeted as a biomarker in the field-effect transistor (FET) based biosensing device reported by Seo et al. [70], where the authors used antibody biofunctionalized graphene sheets as sensing elements achieving LOD of 2.24 x10² copies/mL in clinical samples (Fig. 4(f)).

Other semiconducting nanostructures such as Si, CdSe, and lanthanide materials have also been reported to detect SARS-CoV-2 proteins or antibodies [72-76]. Shan et al. [74] reported the label-free detection of volatile organic compounds (VOCs) in exhaled breath of patients using nanostructured platinum-silicon electrodes to distinguish those affected by COVID-19 from other common lung conditions (Fig. 4(g)). Due to the rapid response time of the device, it can be used for testing large populations if an adequate sterilization process is added to the platform. Sensitive electro-transducing materials such as In₃O₅ nanowires (NWs) could also be surface engineered to detect the N protein of SARS-

### Table 2

| Material                          | Target Virus       | Biological recognition element (antibodies) | Signal response                        | Reference |
|-----------------------------------|--------------------|--------------------------------------------|----------------------------------------|-----------|
| Zinc oxide nanowires (ZnO NWs)    | SARS-CoV-2         | IgG (CR3022)                               | Electrochemical impedance sensing (EIS) | [101]     |
| Patterned zinc oxide nanorod networks | Influenza H1N1 | Anti-H1N1                                   | Cyclic Voltammetry (CV)                | [96]      |
| CdSe/CdS/ZnS quantum dot (QD)     | Influenza H1N1 and H3N2 | Anti-influenza A 7307 monoclonal           | Fluorescence                            | [97]      |
| Graphene/Zinc oxide nanocomposite | Influenza H5      | Anti-fluorescein antibody                   | Cyclic voltammetry (CV) and amperometry | [98]      |
| ZnO NRs in Polydimethylsiloxane (PDMS) | Influenza H1N1, H5N1 and H7N9 | H1N1, H5N1, and H7N9 antibodies             | Electrical response                     | [99]      |
| CdSe/ZnS Quantum dot nanobeads (QDNBs) | Influenza A/H1N1 | Anti-influenza virus A/H1N1                  | Fluorescence                            | [100]     |
|                                   |                    | Anti-influenza A (H7N9) Monoclonal          | Fluorescence                            | [272]     |
LFIAs tests have also been developed using lanthanide-doped nanoparticles by Chen et al. [76]. The sensor allows the detection of IgG antibodies against SARS-CoV-2 in human serum samples with a response time of about 10 min.

In addition to the optoelectronic biosensor response, some authors have reported the use of magnetic nanoparticles (MNPs) (e.g., iron oxide nanoparticles, zinc ferrite) to enhance the RNA extraction process for boosting conventional PCR test efficiency.
or to detect the specific sequences of SARS-CoV-2 [14,21,78]. Therefore, this kind of system offers a practical alternative to the traditional nucleic material extraction process as well as sensitive and specific optomagnetic based sensors that use the response of the MNPs to detect the presence of the virus analyte in the femtomolar range [79]. In the case of Mexico, until now, more than 20 investigation projects to confront the COVID-19 pandemic have been developed by different National Universities and Research Institutes, where the role of magnetic hybridization biosensing probes and fluorescence microfluidic platforms stand out as cost-effective and time-reliable alternatives [80-85]. Additionally, Rodriguez-Moncayo et al. [85] reported the performance of a fluorescence microfluidic platform to detect IgG and IgM antibodies against spike (RBD, and S1 subunit proteins) and nucleoplasmic, showing a specificity in the range of 90–100%. The device is made with PDMS (polydimethylsiloxane) and has a response time of approximately 2 h, and it can be considered mutually cost-effective and time reliable. These characteristics highlighted the biosensor as both a selective and mass scalable opportunity. The state of the art of different nanobiosensing platforms is shown in Table 1 [12,21,23,63,65,67,70,71,74–76,79].

Nevertheless, ZnO nanostructures are highlighted as promising biosensing candidates due to their structural, optical, and electric properties [86-88]. These materials could be surface engineered for the detection of the SARS-CoV-2 virus, and as mentioned earlier, evidence of this is the pending patent of Thevasahayam et al. [23]. This work presents the application of ZnO NRs for the electrochemical detection of SARS-CoV-2 spike protein by an antibody-based approximation. However, developing a biosensing platform to tackle the current global health emergency through the available nanomaterials and different bioselective strategies remains an ongoing challenge.

b. Zn as a suitable tool for detection of emerging respiratory viruses: Challenges and perspectives

As stated before, different materials are studied to establish their sensing performance towards the emerging respiratory viruses (SARS, Influenza, SARS-CoV-2). Hence, the need for the development of complementary techniques to the existing traditional methods is constantly increasing. The actual pandemic situation around the world has shown the lack of infrastructure and trained personnel to solve the demand for tests; that represents a window of opportunity for developing biosensing devices.

Biosensors can be divided into various groups according to their transduction system, such as optical, piezoelectric, and electrochemical [89]. F. Narita et al. [90] reported different materials that can be used as piezoelectric biosensors (Fig. 5 (a–c)) for the detection of human papilloma, vaccinia, dengue, Ebola, influenza A, human immunodeficiency, and hepatitis B viruses. Diverse types of used strategies for pathogenic detection are based on the antibody-antigen reaction; as a result, different immobilization processes have been developed on materials such as gold electrodes, platinum, and graphite, as well as the use of highly crystalline materials, is promoted [91]. However, these materials are expensive, and their processing at a large scale is a challenge for the generation of single-use devices. Therefore, it is desirable to use low-cost and biocompatible materials with a sensitive response signal. In this sense, the platforms based on Zn are a viable alternative to overcome these limitations. Additionally, implementing plasmonic ZnO-based nanocomposites for biological detection could reduce operational costs without losing the optical advantages offered by the material, and the same benefits can also be fetched by utilizing biosensors based on Surface Enhanced Raman Spectroscopy (SERS) substrates.
Numerous researchers worldwide are currently working on ZnO biosensing maturing projects. In our group, different approaches have been studied recently to observe the influence of morphological properties of these nanostructures for various applications, including bio and gas sensors [87,88,92,93] catalysis [94], among others. A. Galdamez et al. [87] reported the orientations of the Au orchestrated ZnO nanowires (ZnO/Au NWs) on the detection sensitivity of DNA using optical and morphological studies (Fig. 5 (d & e)). Due to the knowledge generated from detecting different viruses, Zn-based platforms have an important future projection [95].

In the area of detection of respiratory viruses, Jang et al. [96] reported the fabrication of an immunosensor based on patterned ZnO nanorods (NRs) networks for influenza (H1N1 SIV) detection. Although an Au electrode was used in the device, ZnO played a fundamental role in the immobilization process of the captured antibody. Nguyen et al. [97] developed a CdSe/CdS/ZnS quantum dot (QD) fluorescent dye to detect H1N1 and H3N2 influenza A virus. Even though it is not a complete Zn-based system, the use of this material is essential for bioconjugation (discussed earlier), which allows the detection of the target analyte (Fig. 6(a)). Low et al. [98] reported an electrochemical sensor platform for detecting Avian Influenza H5 (H5N1) based on graphene/ZnO nanocomposite. This work has an interesting approach, as the results obtained from the amperometric study were compared with the efficiency of a conventional agarose gel electrophoresis, finally suggesting a robust validation for the use of these types of biosensors. Further, Ji-Hoon Han et al. [99] reported a nano-flow immunosensor based on ZnO NRs grown on a PDMS sensor for H1N1, H5N1, and H7N9 influenza virus detection using an electrochemical method. The ZnO NRs allowed the immobilization of different antibodies, as well as an increase insensitivity. Nasrin et al. [100] employed ZnO as a precursor in the development of fluorescent CdZnSeS/ZnSeS quantum dots (QDs) and gold nanoparticles (AuNPs) for the detection of influenza virus A/H1N1 (Fig. 6(b)).

Table 2 presents a summary of the works involving the use of Zn-based nanomaterials to detect emerging respiratory virus diseases.

From these works, a retrospective is opened about Zn-based nanomaterials and their potential use in SARS-CoV-2 detection. These promising devices could develop trials to involve specific antibodies against the well-known spike protein S1. Xiao Li et al. [101] reported an experimental approach to enhance the biosensing performance of paper-based electrochemical impedance sensing (EIS) nanobiosensors with working electrodes (WEs)
decorated with vertically grown ZnO NWs. These nanobiosensors can differentiate the concentrations (blank, 10 ng ml⁻¹, 100 ng ml⁻¹, and 1 μg ml⁻¹) of IgG antibody (CR3022) to SARS-CoV-2 in human serum samples (Fig. 5(f)). Furthermore, Thevasahayam et al. [23], also reported the rapid detection of SARS-CoV-2 by employing ACE2 functionalized ZnO NWs.

As discussed, to detect different respiratory viruses, Zn presents a wide range of characteristics in the various platforms. In some cases, as a part of a nanocomposite or in other cases, it is even used in the union of functional groups to recognize biomolecules (Fig. 7(a)). Hence, it is possible to take advantage of signal transduction strategies to develop different types of optical or electrochemical biosensors. The development of immunosensors is based on the specificity of the antigen–antibody reaction. COVID-19 disease is reported to induce acute antibody (IgM, IgG) response, and the levels of antiviral antibodies are plateaued within 6-days after seroconversion [102]. The spike protein S1 protein is one of the main references for COVID-19 detection. This could function as an indicator of the virus even in the incubation period and for asymptomatic cases [103-105]. Nevertheless, it is suggested that other envelope or membrane proteins can be used to identify different SARS-CoV-2 variants. This could allow the generation of more selective biosensors based on the knowledge, response signals, and characterized behavior of devices based on nano Zn and related compounds. For this reason, ZnO can also be used as a highly sensitive platform for the immobilization of antibodies;
consequently, a “serology” test can be performed to measure antiviral antibodies in the blood [106]. In addition, the current knowledge about nano Zn handling and its advances for detecting low analyte concentrations could reduce the analysis times in different population sectors.

According to the previous explanations, some of the functionalization strategies based on silane group immobilization can be employed suitably to use merely nano Zn and its compounds as a prospective biosensor platform. ZnO is one of the materials used for biofunctionalization [88]; since it is an oxide, the generation of OH\(^-\) groups is favored compared to other metallic materials that sometimes require more complex treatments. Hence, in this case, silanization represents an advantage in performing functionalization strategies, resulting in more OH\(^-\) groups available for binding.

Fig. 6. (a) Schematic view of a fluorescent test showing the application of the conjugate and sample to rapid diagnostic strips. Copyrights Elsevier 2020 [97]. (b) Diagram for the preparation of CdZnSeS/ZnSeS QD-peptide-AuNP nanocomposite and influenza virus detection. Copyrights Elsevier 2020 [100].

with the biological recognition element. For instance, Allen et al. [107] showed the analysis of treatments using triethoxysilane on ZnO thin sol–gel grown films, emphasizing the role of silane in this material. Corso et al. [108] reported the use of two (3-glycidyloxy propyl)-trimethoxysilane (GPS) and (3-mercaptopropyl)-triethoxysilane (MTS) for the immobilization of IgG antibodies on ZnO surfaces.

Moreover, García et al. [109] evaluated the effects of silanization with amino-propyldiethoxymethylsilane (APDEMS) on hydroxylated sidewalls of ZnO-NWs, highlighting that some studies only focus on the binding of a biomolecule; however, just a few studies address the effective surface functionalization of ZnO. The reported functionalization strategies show a proposed method for their adaptation on Zn-based biosensing devices (Fig. 7(b)), which leads to the chemical modification of a surface for the antibody immobilization [93] and an optical transduction response. Therefore, it represents an affordable alternative for research groups that have developed biosensors using silanes that could be adapted to immobilising recognition agents against the spike S1 protein (SARS-CoV-2).

In addition, it is noteworthy to mention that the preparation of biosensors faces diverse challenges and offers a comprehensive perspective and versatility in the assembly. For example, the strategies for antibody immobilization using crosslinkers improve the orientation in different materials giving a better sensibility; however, it raises the overall cost. Another challenge is the coupling of response signals of biosensors with electronic and portable devices that are subjected to international quality standards for daily use. Lastly, the validation of detection limits and its comparison with molecular reference methods is still an issue to be further solved. On the other hand, random immobilization strategies can also affect the sensitivity of these devices. The increasing knowledge in this health emergency proposes the investigation of improved sensor surfaces from different nanomaterials such as Zn and its compounds to achieve fast detection response and high selectivity and sensitivity. Already gained information with Zn-based biosensing platforms for other respiratory viruses could be a surplus and could act as a reference guide to design these new potential biosensors for early detection of diseases. According to the databases consulted, there is a limited number of works related to the detection of COVID-19 with nano Zn and its compounds, which opens the panorama for future research and studies.

3. Antimicrobial activity of Zn-based materials for COVID-19 disinfection

Apart from the above-mentioned biosensing platforms, another essential method to constraint COVID-19 is the development of self-sterilizing materials. Up to this moment, it has been accepted
that the primary source of transmission of SARS-CoV-2 as well as other respiratory viruses occurs through liquid droplets and aerosols coming from the exhalation or sneeze of infected persons [110]. While the use of personal protective equipment (PPE) such as face masks considerably reduces the probability of contagion, continuous and prolonged use of this type of equipment requires meticulous subsequent disinfection processes to assure reusability and/or responsible disposal [111].

Likewise, public space facilities such as hospitals, schools, and transports are also recommended to be sanitized regularly as the indirect contagious by fomites could lead to a viral spread in the community [112]. Thus, the development of effective viral disinfectant materials is of great importance, and a complementary approach that can tackle the current pandemic and possible outbreaks of emerging viruses is of high interest (Fig. 8(a)). In this regard, metallic antiviral nanoparticles are the most reported and studied systems due to their inherent broad range of antimicrobial activities and effectiveness at a much lower dosage than their bulk counterparts [10,113]. Nevertheless, Zn-based nanomaterials have proven to be efficient antimicrobial materials that offer diverse valuable photocatalytic, surface, and morphological properties to inhibit and deactivate pathogens. In other words, ZnO nanostructures exhibit tunable antibacterial, antifungal, and antiviral capacities.

In addition, cost-effectiveness and low cytotoxicity to various human cell lines have led to the practical implementation of nanostructured Zn-based materials into food packaging applications to maintain the proper hygiene and prolong the quality of intake products [27,114,115]. ZnO NPs have also been implemented in the development of water purification systems [116]. In the work of Munnawar et al. [117], the obtention of ZnO NPs loaded with chitosan and their cytotoxic activity towards fungi and bacteria is presented. Even when chitosan biopolymers exhibit inherent antibacterial properties, the hydrophilicity and porosity of the membrane were further improved by incorporating ZnO NPs. The authors reported a population reduction in the treated samples with a 15% loading of ZnO NPs. Thus, the synergic effect between organic matrices and inorganic reinforcement phases can be further explored for more disinfectant applications, as is also pointed out by Mizielinska et al. [115].

The properties mentioned above illustrate the cytotoxic mechanisms that ZnO exhibits on various cellular components, and it has shown high effectiveness against a wide variety of Gram-positive (Staphylococcus epidermidis, Streptococcus pneumoniae, Staphylococcus aureus), Gram-negative (Klebsiella pneumonia, Campylobacter jejuni, Escherichia coli) bacteria and fungi (Candida albicans, Aspergillus Niger) [118-123]. The antimicrobial and antifungal mechanisms are strongly influenced by parameters such as the nanostructures concentration, size, morphology, and surface functionalization [27,124], which can be precisely controlled with the different deposition techniques. For instance, Cha et al. [125] described the effect of different morphologies of ZnO nanostructures (spherical and pyramidal) over the antibacterial and enzyme inactivation activity. The authors reported a biomimetic design of pyramidal NPs to act against super-resistant bacteria (methicillin-resistant Staphylococcus aureus), which is presented in most hospitals. Another experiment comparing the antibacterial activity of different morphologies of ZnO nanostructures is presented in the work of Lopez et al. [126]. In this case, the authors observed an enhancement in the microbial inhibition of E. coli and S. aureus by ZnO nanotubes (ZnO NTs) in comparison to the one displayed by commercial ZnO NPs. The ZnO NTs morphology aids in an increase in the surface area and also reduces the aggregation in the system, thus favoring antibacterial activity.

Even though the precise mechanisms of the antibacterial/antifungal activity of ZnO nanostructures remain under debate, cellular membrane-nanoparticle electrostatic interaction, reactive oxygen species (ROS) generation [123,127,128], and intracellular Zn²⁺ release [129,130] are expected to be responsible for reducing cell viability (Fig. 8(b)). As pointed out by Sirelkhhatim et al. [27], the generation of H₂O₂, O₂⁻ and *OH ROS species by ZnO nanomaterials plays a significant role in localized cell wall damage by oxidative stress, in some cases leading to cell death. Other proposed mechanisms include the penetration of neutral ROS into the cell as the cellular membrane repelling the negative species. Then the chemical oxidation process could interrupt the respiratory activity of the cell, thereby producing DNA damage or protein denaturation [114,131] (Fig. 8(c)). A comparative study of the ROS generation for bacteria and/or fungal control applications of several metal oxides [132]. They compared the performance of ZnO with other typ-
ical binary semiconductors, corroborating the potential and relevance of the material in this research field. In the current COVID-19 pandemic, the antibacterial properties of ZnO materials could be targeted towards opportunistic diseases linked with SARS-CoV-2 infection, such as pneumonia (caused by bacterial or fungal pathogens) which is responsible in some cases for fatal complications in the patient [133].

Although most of the publications found in the literature are focused on the antibacterial/antifungal properties of nanostructured ZnO materials, some authors also report the application of these nano-systems as antiviral agents against viruses such as Herpes simplex virus (HSV-1), H1N1 influenza virus, and recently against the SARS-CoV-2 virus.

Among the first works to study the effect of nanostructured ZnO as an agent to reduce the infectious capacity of a virus is the one reported by Mishra et al. [134]. The authors reported the obtention of ZnO’s micro-nano structures (MNSs) with a morphology that bio-mimics the cellular membrane receptors that the HSV-1 virus typically targets. The viral inhibition mechanism is based on the electrostatic interaction between the negatively charged surface of the ZnO nanostructures and the positively charged protein surface of the virus. As a result, the ZnO MNSs are found to function as anchor points for HSV-1 virus particles and thus preventing them from reaching the target cell gD-receptors. The former mechanism could be used to develop ZnO nanostructures that target SARS-CoV-2 surface proteins (Fig. 9(a)).

In the work of Tavakoli et al. [135], a more recent study about the antiviral properties of ZnO NPs against the HSV-1 virus is presented. Herein, a surface functionalization approach using polyethylene glycol (PEG) was developed to increase the antiviral response and reduce the cytotoxicity of the NPs. A similar ZnO-PEG functionalized NPs approach is reported by Ghaffari et al. to perform against the H1N1 influenza virus [136]. Cytotoxic analysis revealed that implementing a PEG layer significantly enhances the biocompatibility of ZnO NPs towards Madin-Darby canine kidney MDCK culture cells. The smaller the size of NPs, efficient is the

Fig. 9. Possible mechanisms of antiviral activity of ZnO against SARS-CoV-2. (a) Design of ZnO nanostructures for the possible anchoring of SARS-CoV-2 virons, thus inhibiting interaction with host cell receptors. (b) Internalization of ZnO nanostructures for the inhibition of early stages of the viral replication cycle. (c) Ion release as a surface attack mechanism to disrupt the plasmid and RNA virus integrity. (d) Photocatalytic generation of reactive oxygen species for the possible degradation of the lipid, protein, and nucleic structure of SARS-CoV-2. Figure created with BioRender.com.
cellular absorption. The authors propose that the antiviral activity of the material is carried out when the NPs are internalized in the host cells (Fig. 9(b)). Some recent publications have been made over the direct antiviral properties of ZnO nanomaterials against the current SARS-CoV-2 virus. In the work of El-Megharbel et al. [137] According to the proposed mechanism, ZnO NPs are tested towards Vero-E6 cells infected with SARS-CoV-2, resulting in the virus inactivation by Zn²⁺ release and ROS formation (Fig. 9(c)). The authors pointed out that the same mechanisms for antibacterial activity are also responsible for damaging the lipid membrane and RNA and thereby inactivating the virus. However, the cytotoxic studies reflect healthy cell apoptosis at relatively low concentration levels of ZnO NPs, suggesting the need to improve their biocompatibility.

Furthermore, another molecular docking analysis about the interaction between the adhesion proteins of the SARS-CoV-2 virus and ZnO nanostructures is presented by Adhikari et al. [138]. In this work, the authors determined that the interaction between the spike protein of SARS-CoV-2 and the ACE-2 receptor is comparable in energy values, with that experienced between the spike adhesion protein of the virus and the surface of both 6D spherical NPs and the (1 0 0), (1 0 1) and (0 0 2) crystal faces of 2D ZnO nanostructures. The computational study also reflects the viral denaturalization posterior to the adsorption in the semiconducting surface. This feature makes the material a suitable option to be applied in face mask manufacturing. This application was also explored by Adhikari et al. when manufacturing ZnO nanoflowers (NFs) on cotton fabrics substrates using a low-temperature hydrothermal approach. The implementation of antiviral organic compounds has previously been reported; however, the thermal and chemical stability of the latter is still among the main challenges to be overcome. This is one of the reasons why obtaining antimicrobial materials at low temperatures is of great interest for their incorporation into fabrics. As a proof of concept, the authors studied the antimicrobial activity against the model bacteria Pseudomonas aeruginosa. The pathogen was targeted due to its LecA membrane protein which confers its mimicry with the spike protein of SARS-CoV-2. The results showed an antibacterial effect through the rupture of the cell membrane, a behavior that was maintained even after washing the fabric 50 times. Scanning electron microscopy (SEM) images also showed minimal fiber deterioration even after several washes. This suggests the effective implementation of functionalized fabrics with ZnO nanostructures in washable antimicrobial clothing and personal protective equipment.

Furthermore, the photocatalytic properties of ZnO nanostructures can also be used to develop photoinduced self-cleaning coatings, where good adhesion to the substrate, improved broadband photoabsorption, and innocuous biodegradability of the photocatalytic platforms are among the main goals to be achieved [55]. In fact, during the health emergency of COVID-19, street clothing and personal protective equipment. This suggests the effective implementation of functionalized fabrics with ZnO nanostructures in washable antimicrobial clothing and personal protective equipment.

4. Zn as an essential element in emerging virus mediated-respiratory diseases

It has been found that the emergent lethal respiratory viruses acquire newer survival tactics to live inside host cells and trick the host's immune system. For instance, novel coronavirus and drug-resistant species of influenza viruses call for an urgent requirement of novel treatment strategies [140,141]. Antiviral agents directly act on the virus or intracellular paths for viral replication by several mechanisms such as attacking target proteins, acting as ligand analogs, and disturbing the viral replication cycle. Nevertheless, most available antiviral agents exhibit disadvantages, such as low solubility, minor stability, high potential of drug-drug interaction, and high toxicity, which should be further improved. Therefore, this clearly remarks the need for further research in this respective area [7,140]. Nanomedicine has come to be one of the most promising technologies because of its ability to combat viral infections facing the restrictions of conventional therapies along with nanoformulations, including solid lipid NPs, polymeric NPs, liposomes, dendrimers, micelles, self-assembled nanoemulsions, and cyclodextrins [7,56]. Metallic NPs exhibit inherent antiviral activity and are extensively studied for this purpose; among them, Zn has been found efficacious against respiratory infection-causing viruses [7]. The role of Zn-associated compounds, Zn conjugates, and Zn ionophores have been studied against respiratory viruses [142-144]. In the present COVID-19 pandemic situation, specific treatment for this infection has not yet been determined; however, the therapeutic options are mainly aimed at improving respiratory function by Oxygenation and Ventilation and controlling the advanced symptoms through diverse Pharmacologic Interventions [145]. In this context, medical professionals are starting to consider the responses through supplements such as vitamin D, vitamin B12, probiotics, and Zn [146-148]. Likewise, clinical trials have been performed to determine the effect of Zn supplementation on virus-mediated-respiratory diseases, which will be analyzed below, along with the potential relevance of their nanostructured form, toxicity, and role as a supportive treatment and prophylaxis of emerging respiratory infections.

a. Zn supplementation as a supporting treatment

Zn is a vital mineral, and it regulates numerous enzymes and proteins that participate in necessary cellular events, mostly related to inflammatory system regulation [147,149]. Zn in its bulk form is registered safe by the United States Food and Drug Administration (FDA) [21 CFR182.8991]. The standard route to obtain this element is through the ingestion of enriched food or nutritional supplements, while the Recommended Daily Intake (RDI) values are in between 3 mg and 16 mg; having an oral Lethal Dose (LD50) close to 3 g/kg per body weight [150]. The main Zn compounds used for supplements are zinc gluconate (C₁₂H₂₂O₁₄Zn), zinc citrate (C₁₂H₁₁O₇Zn), zinc sulfate (ZnSO₄), bis-glycinate (C₄H₈N₂O₄Zn), zinc acetate (C₂H₄O₂Zn), zinc picolinate (C₇H₇N₃O₅Zn), zinc oxide (ZnO),
zinc chloride (ZnCl$_2$), and zinc aspartate (C$_8$H$_{12}$N$_2$O$_8$Zn), among others [151,152]. Depending on the Zn compound, its functionality could vary for their use as supplementation; for instance, zinc acetate (C$_4$H$_6$O$_4$Zn) has been more effective than zinc gluconate (C$_{12}$H$_{22}$O$_{14}$Zn), and lastly, zinc citrate (C$_{12}$H$_{22}$O$_{14}$Zn$_3$) for the common cold [153]. Notably, it has been found that zinc gluconate (C$_{12}$H$_{22}$O$_{14}$Zn), zinc citrate (C$_{12}$H$_{22}$O$_{14}$Zn$_3$), and zinc sulfate (ZnSO$_4$) are absorbed better in healthy people, while ZnO is least absorbed.
Zn is crucial in regulating inflammatory cytokines by reducing IL-15 [154,155]. However, in a different study, it was found that along with additional food supplementation, ZnO absorption was equivalent to that of ZnSO4 [156]. Furthermore, Zn can also be used as a food fortifier, and in that case, bioavailability will be affected by interactions with different food components or through proper process handling. Therefore, it is essential to study the compound solubility, charge density, reduction potential, pH, complex formation, and processing impact before manufacturing [156]. However, Zn performance depends on its ion availability from different salts, for example, zinc chloride (ZnCl2), ZnSO4, and zinc acetate (Zn(OAc)2), as well as the target virus [157].

On the other hand, previously reported studies established that nearly two billion people in the world may have prolonged Zn deficiency, thus showing related clinical problems like growth retardation, cognitive impairment, liver, and renal diseases, chronic inflammation, and triggering oxidative stress, among others [158]. The studies have also demonstrated that plasma Zn levels are markedly reduced through infections, and there is a correlation of around 16% of the world’s severe respiratory infections and bacterial or fungal co-infections [159,160,161,151]. A representation of around 16% of the world’s severe respiratory infections and bacterial or fungal co-infections [159,160,161,151].

Table 3

| Type of intervention of Zinc against SARS-CoV infection | Suggested Antiviral mechanism | Reference |
|---------------------------------------------------------|-------------------------------|-----------|
| Zn inhibits RNA-dependent RNA polymerase (RdRp) of SARS-CoV | Decreases the rate of virus transcription | [52] |
| Zn may induce inhibition of S. pneumoniae growth | Modulation of confection of bacterial Mn (II) dependent homeostasis | [188] |
| Zn binds to the SARS-COV enzyme catalytic residues | Inhibition of SARS-CoV activity | [53,158] |
| Zinc may lower plasma C-Reactive Protein (CRP) concentration | Downregulating the response to inflammation via CRP | [161] |
| Zn exposure may decrease recombinant human ACE-2 activity and expression | Modulating effect on SARS-CoV-2/AACE2 interaction | [189,190] |
| Regulate tight junction proteins ZO-1 and claudin-1 | Controlling Antioxidant and Anti-inflammatory activity of respiratory epithelium | [171,172] |
| Control of IFNα production and intensifying its antiviral activity through IFNα-induced JAK1/STAT1 signaling | Increasing antiviral activity of cytokines | [173,174] |
| Inhibition of caspases-3, 6, and 9, and an increase of the Bcl-2/Bax ratio expression | Enhances cells resistance to apoptosis | [175] |
| Zn exerts its anti-inflammatory activity through inhibition of IRK activity and subsequent NF-kB and modulation of regulatory T-cell functions | Down-regulation of pro-inflammatory cytokine production and increases the cytotoxicity of NK and T cells | [167,176,179] |
| Inhibition of ADAM 17 enzyme at the Zn cofactor site inactivates the enzyme, causing downregulation of inflammation | Downregulation inflammation via of ADAM enzyme | [180] |
| Zn supplementation improves mucociliary clearance, aid with the removal of bacteria and viruses and, diminishes inflammatory reaction and lung damage | Improving defense of the airways and bronchial epithelium | [151,170,171] |
| Zn can counteract virus fusion with the host membrane, impairs protein translation and processing, blocks viral particle release, and destabilize the viral envelope. | Improving effects on host cell metabolism to prevent damage from infection | [153,183] |
| Zn supplementation can reverse lymphoplasia, Attenuated lipopolysaccharide-induced hyperactivation, recruitment, and formation of neutrophil extracellular traps | Improving Immune system cells response | [182,183] |
| Zn supplementation might reduce the harmful microbes and increase the helpful ones in a dose-dependent manner | Prophylaxis based on gut microbiota profile to improve immunity | [184-187] |

Furthermore, Zn homeostasis is crucial for wound healing and tissue recovery when it is linked to inflammation or mechanical damage. Also, Zn has been associated with ROS production in platelets, indicating a decrease in thrombus formation and inhibiting other complications by respiratory virus infections [151]. Furthermore, it has been studied that Zn affects the IFN-α binding to IFN receptor 1 (Lambda interferons), ensuing in reduced antiviral activity (Influenza virus) in vitro, in which the mechanism of Zn-mediated inhibition could occur in an extracellular manner through degradation of viral RNA, and thereby abrogate viral infectivity [168]. Consequently, there is a suggestion to monitor Zn levels in the high-risk population (elderly and patients with comorbidities) to achieve the most beneficial therapeutic value [146,169,151]. Furthermore, Fig. 10(d) exemplified Zn’s in vitro evidence of antiviral properties in emergent respiratory virus infections [165]. State-of-art in vitro evidence of possible Zn mechanisms for antiviral action by supplementation is described in Table 3 [52,53,151,157,158,161,167,170-190].

Moreover, it can be suggested that Zn use as a supplement may confer various benefits associated with treating emerging respiratory virus infection, considering that its use is safe at controlled doses, has a positive role in inflammatory response modulation and antiviral activity. Likewise, it has been revealed that Zn treatment diminished the length of common cold signs in healthy persons [191], can restrict influenza virus infections [153], had antiviral action in rhinovirus and respiratory syncytial virus (RSV) infection, as well as influenza-MRSA bacterial superinfection [53,192]. According to the COVID-19 Treatment Guidelines of the National Institute of Health U.S.A. [193], there is still insufficient data to recommend either for or against the use of Zn to treat respiratory infection viruses. However, few studies were recently registered worldwide in the National Library of Medicine of the United States and in the European Union Clinical Trials to assess Zn efficacy in treatment combined with drugs or prophylaxis against complications of SARS-CoV-2 and Influenza (flu) infection [194,195]. In the analyzed databases, the use of ZnSO4 and C12H22O14Zn in concentrations between 50 and 220 mg can be understood (Fig. 10(e & f)). Nevertheless, no conclusive results have been reported so far, and this is a topic of keen attention among...
different research groups [196-205]. It should be highlighted that the outcomes that can be modified in virus infection with the use of Zn are too diverse and require a separate analysis in each case (from the reduction of infections, hospitalizations, and intubations to the decrease in deaths). Also, it is crucial to evaluate the effect of Zn supplementation in infected patients and to identify if they have a preceding deficiency of this mineral or not before initiating treatment.

Moreover, many investigators recently have pointed out the importance of nanomaterial-based technological resolutions to combat emerging respiratory virus infections [206,207]. As revealed, nanomaterials as supplements have a better capacity than conventional sources due to their form, easier uptake by the GI tract, and enhanced effectiveness at lower doses [208]. The following section discusses the state-of-the-art of Zn NPs studies against the emerging respiratory viruses.
b. Potential of nano-Zn compounds against respiratory viruses

This recent coronavirus pandemic has shown promising results, seeking complementary therapy to support prevention, treatment, and recovery. In this context, a few studies have investigated the properties of metal NPs to augment their selectivity and efficacy against respiratory viruses since these NPs can bind with the viral proteins (via Van der Waals forces) and can help to conduce its inactivation \cite{7,209}. Correspondingly, in double-stranded RNA viruses, Ag NPs have shown that after interaction with the viral genome, they inhibit viral replication; likewise, Cu NPs inhibit virus-cell binding and attachment and destroy the viral genome. Whereas Fe NPs bind to the virus to prevent it from attaching to the host cells. Though, Se NPs guard from apoptosis caused by the infection of the virus \cite{210}. In the case of Zn NPs, it has been observed that it interferes with viral DNA polymerase activity and binds to viros resulting in inhibition in viral replication and entry \cite{7,210,211}. Although the exact mechanism for Zn nanomaterials to act as agents for the treatment of the recurrent respiratory virus is still under investigation, there is evidence showing that the performance of Zn depends on the ion availability from different composites. It further depends on NPs morphological aspects such as particle size, shape, concentration, agglomeration, colloidal formulation, and media pH \cite{7,209}. In this context, PEGylated ZnO NPs have better antiviral activity than bare ZnO NPs, significantly reducing H1N1 virus titer post-infection \cite{135,136,158}. Likewise, the antiviral activity of ZnO-PEG-NPs is dose-dependent compared with an antiviral drug, noticing the greatest H1N1 antiviral influence at 75 μg/mL with an inhibition rate of 52.2% \cite{11(a)} \cite{136}. In antiviral therapy, ZnO NPs offer advantages and attempt to deflect the occurrence of drug resistance. In this context, an investigation employing ZnSO₄ (1.5 mg/mL) and Ag NPs showed comparable potentiated antiviral action with epigallocatechin gallate (EGCG) (50 μM) against the H5N1 avian flu virus in embryonated SPF eggs, further helping to limit the infection resistance and without been affected by virus mutations \cite{11(b)} \cite{212}. Another study with a composite of Au/Ag/ZnO-NP (TPNT1) showed inhibition of different strains of SARS-CoV-2, human H1N1, and avian H5N1 influenza viruses, including the oseltamivir-resistant strains, proposing it as a prophylactic agent against SARS-CoV-2 and opportunistic infections by oral gargling, nasal spray, nebulized inhalation or even systemic use after an appropriate clinical trial \cite{11(c)} \cite{141}. Moreover, docking molecular studies have been performed to determine the interaction between ZnO NPs and the common SARS-CoV-2 targets. One of the studies speculated the possible interaction among ZnO NPs and ACE2 receptor, COVID-19 RNA-dependent RNA polymerase, and the main protease, putting forward the attention towards the binding of ZnO NPs with the tested targets and observing an enhanced dose-dependent cellular uptake \cite{11(d)} \cite{158}. In the work performed by Hamdi et al. \cite{214} it was found that ZnO could mimic the inhibition behavior of remdesivir and saquinavir drugs by hydrogen bond interactions with the Arg555, Ser759 amino acid and U10, U20 uracil bases, respectively. The role of hydrogen bond and other interactions as π-cation and π-π are essential in the ligand–protein stabilization indicating the ZnO NPs as a possible treatment to COVID-19. Furthermore, Fig. 12 exemplified the in vitro evidence of Zn NPs antiviral properties in emergent respiratory virus infections.

It should be noted the aforementioned respiratory viruses, even despite using different cellular pathways, share the existence of the ACE2 protein at the beginning of the digestive system and upper respiratory tract (discussed earlier) \cite{215}, which represents an excellent motivation for the proposed therapy with Zn NPs. The results of nano Zn compounds in various models studies for antivi-
antiviral activity by ZnNPs against respiratory virus. 

Can ultimately experience an acute condition of ARDS-like [227].

and/or ZnO, mostly from industrial processes or military activities, 

Table 4 [136,141,158,212,213,216-218].

Table 4
Antiviral activity by ZnNPs against respiratory virus.

| Nano Zn Compound | Species/Study | Effects | Remarks | Ref |
|------------------|---------------|---------|---------|-----|
| ZnO-NPs and PEGylated ZnO NPs | Influenza A/Puerto Rico/8/34 (H1N1; PR8) propagated in Madin-Darby canine kidney (MDCK)-SIAT1 cells and PRV virus infection. | PEGylated and unPEGylated ZnO-NPs led to inhibition rates of 94.6% and 52.2%, respectively of H1N1 virus. | A proposal as an antiviral agent against H1N1 influenza virus infection | [136] |
| Zn-based physiometa-composites (PMCs); Mn, Fe, Ni and Co-doped, ZnO, ZnS or ZnSe | Influenza A Virus (H9N2) in embryonated Specific-Pathogen-Free hen’s eggs | | | |
| ZnSO4 (zinc II) NPs | | | | |
| ZnO NPs | BALB/c mice injected with M. pneumoniae | Reduced total protein, inflammatory cells, inflammatory cytokines (IL-1, IL-6, IL-8, TNF-α and TGF). ZnO/BER complex inhibits spike protein binding with angiotensin-converting enzyme II (ACE II), PL pro activity, spike protein and E protein levels, and expression of both E-gene and RNA dependent RNA polymerase (RdRp). | Potentially antiviral activity of EGCG by co-administering it with zinc II and AgNPs | [212] |
| ZnO Berberine (BER) complex | VeroE6 toxicity, anti-COVID-19 activity | | | |
| Au-NP/Ag-NP/ZnO-NP and ClO2 nanocomposite (TPNT1) | SARS-CoV-2 strains in vitro cell-based assay | Block viral entry by inhibiting the binding of spike proteins to ACE2 receptor and to interfere with the syncytium formation | Effective concentration within the range to be used as food additives provide prophylactic effects against both SARS-CoV-2 and SARS-CoV-1 infections | [273] |
| Au-NP/Ag-NP/ZnO-NP and ClO2 nanocomposite (TPNT1) | (H1N1) and avian influenza A virus (H5N1) cell-based assays | Reduced the cytopathic effects induced by human H1N1 and avian H5N1 influenza viruses, including oseltamivir-resistant virus isolates | Inhibit or prevent viral infection and opportunistic infections | [273] |
| ZnO NPs | COVID-19 targets include the ACE2 receptor, COVID-19 RNA-dependent RNA polymerase in humans in lung fibroblast cells and in silico molecular docking | The binding of ZnO NPs with the three tested COVID-19 targets via hydrogen bond formation was detected. | Infer further biological and therapeutic studies | [158] |
| EGCG-AgNPs/ZnSO4 | Influenza A Virus (H9N2) in embryonated Specific-Pathogen-Free hen’s eggs | Co-treatment with ZnSO4 (1.3 mg/mL) increased the EGCG antiviral effect | May prevent virus transmission, inhibit virus replication and inhibit microbial resistance | [218] |

The permissible exposure limit in the workplace is 5 mg/m3 for ZnO [228]. Similarly, chronic Zn toxicity, which occurs more frequently, happens after long-term high intakes of Zn (150 mg to 450 mg per day) [161], and some of the warning signs comprise lethargy, compromised immunity, and health problems associated with obesity [229]. Excessive Zn intake is related to alterations in Cu and Fe homeostasis at the cellular level, leading to anemia and damage to the nervous system [147]. Some reports indicate that the impact of Zn on apoptosis (programmed cell death) is still under discussion. Zn can either be a pro- or anti-apoptotic element and equally, its deprivation and excess can induce apoptosis. The apoptotic effects are related to regulating the proteins Bcl-2-like, Bax-like, and cytochrome-c in the mitochondria and its intervention in the pathway of Akt/ERK proteins. The anti-apoptotic activities have been recognized by suppressing signaling pathways that regulate apoptosis mainly through caspases and TPEN protein. However, the variables in this system need further research effort [161].

As mentioned above, the leakage of free Zn2+ ions or Zn OH− from ZnO NPs has been suggested to justify their cytotoxicity; nevertheless, as mentioned earlier, some reports indicated that ROS species could be the ones that can trigger their antimicrobial activity [27,114,123,126-129]. Also, it has been observed that ZnO NPs toxicity should be associated with the release of Zn2+ ions rather than with the presence of NPs since they can be easily dissolved in natural aqueous media with a duration expectancy smaller than 90 min, facilitated by the water chemistry, and the presence of dissolved organic matter. Similarly, dissolution rates increased with the addition of strong chelating agents, such as EDTA and L-cysteine. In contrast, a decrease was observed in the...
presence of polymeric organic matter such as sodium alginate [168,230]. ZnO NPs toxicity is mainly determined based on the direct measurement of released Zn\(^{2+}\) ion concentrations under different environmental conditions such as pH, temperature, ionic strength, and chemical composition [231]. Likewise, it has been shown that there is a specificity of biomolecular interactions of Zn nanocomposites with proteins, observing a change in spectral signature when proteins such as ribonuclease A (RNase A) interact with ZnO NPs. It was found that when RNase A was bound to Mg/ZnO, the intensity was quenched, while it was magnified when taurine yeast RNA was bound to RNase A and Mg/Zn [232].

In a human inhalation study, it was found that healthy adults inhaled approximately 500 g/m\(^3\) of ultrafine ZnO, fine ZnO, and filtered air while at rest for 2 h. It has shown no differences between exposure conditions in leukocyte surface markers, hemostasis, and cardiac electrophysiology conducted 24 h post-exposure [233]. In studies in invertebrates have shown that the toxic responses of ZnO NPs are due to their interactions and/or uptake by the blood cells within the insect’s body, and the toxic reactions may diminish once the insects excrete these NPs [234]. In another study that explored the role of the group of proteins that bind to the NPs surface (protein corona) for biokinetic and toxicology propose, they found that the protein corona formed on silica-coated ZnO NPs had higher amounts of plasma proteins, mostly albumin and, transferrin, compared to the NPs without the coat, showing that surface modification with amorphous silica alters the protein corona, agglomerate size, and zeta potential of ZnO NPs, which in turn influences ZnO biokinetic behavior in the circulation. This emphasizes the critical role of the protein corona in the biokinetics and toxicology of ZnO NPs [235]. An investigation described that corona formation occurs in the alveolar lining fluid and comprises plasma proteins, a surface-active phospholipid (PL) protein mixture, and a layer of aqueous hypo phase. In this scenario, ZnO NPs incubated in rat lung lining fluid in vitro binds mainly with albumin, transferrin, and \(\alpha\)-1 antitrypsin. A more extensive database of corona composition of a varied NP library should be developed to help predict the effects and biokinetics of inhaled or ingested NPs [236].

The influence of serum Zn concentrations is the primary health outcome for clinical applications of Zn-based biomaterials. In an earlier study, elevated serum Zn levels were significantly associated with an increase in total spine and femur bone mineral density; also, each 10 µg/dL increase was associated with an increase in the risk of diabetes mellitus, cardiovascular diseases, and coronary heart disease in participants with serum Zn levels \(\geq 100 \mu g/dL\). However, it had no significant associations with the risk of fractures, congestive heart failure, heart attack, thyroid disease, arthritis, osteoarthritis, rheumatoid arthritis, dyslipidemia, and cancer [237]. Furthermore, the main problem that has been reported regarding the interface among Zn NPs and diet components is that it might affect nutrient absorption, could modify the expression of nutrient-related genes, or, as mentioned earlier, could induce oxidative stress [136]. Although, it has been reported that most of the observed toxic effects by ingestion of ZnO NPs are reversible after vitamin C or Quercetin supplementation, among other antioxidants [238,239]. The treatment for Zn toxicity is directed to overcome the symptoms and restore Cu or Fe levels, though severe cases may also require chelation therapy and specialized medical attention [222]. However, discontinuing Zn supplementation may suffice and resolve the symptoms [161,240].

In this regard, toxicity investigations with ZnO NPs, have shown that the oral administration of ZnO had no impact on organ weight, intestinal microbiota, and other mineral concentrations whereas, ZnSO\(_4\) appeared to incite further toxicity [241]. It has been found that ZnO and ZnO NPs at applied concentrations do not affect the growth performance and composition of blood minerals in Mar-khoz goats [242]. Zn-deficient rat models were supplemented orally with nano ZnO fortified Biscuits (13.5 ppm, 27 ppm, and 54 ppm) and were compared with ZnO from the bulk. Results showed an enhancement of body growth rate, appetite, and hair growth, without mortality and neither histopathological abnormalities in the Liver, Kidneys, and Testis. Recommending 13.5 ppm nano Zn fortified biscuits for managing mild Zn deficiency and the dose of 27 ppm for more severe conditions [243]. Likewise, considerable improvements were observed in the health status and birds’ immunity supplemented with nano Zn to broiler diets at 0.06 mg/kg compared with the conventional dose of 15 mg/kg of organic and inorganic Zn with the basal diet [208]. In newly-weaned piglets, replacing high-dose dietary ZnO with low-dose porous and nano ZnO had a similar effect on improving growth performance and intestinal morphology, reducing diarrhea and intestinal inflammation [243]. In a virus mitigation assay, no cytotoxicity of MARC-145 cells was observed for any of the MnZnS NPs concentrations tested (100 µg/ml, 50 µg/ml, 20 µg/ml, 10 µg/ml) [213]. An in vitro pulmonary toxicity test of ZnO NPs in human lung cell lines showed a toxic effect at concentrations between 10 and 50 µg ml\(^{-1}\). Also, 2D cell model submerged exposure revealed toxicity for ZnO at 10 µg ml\(^{-1}\) for hAEC and hAELVi cell lines, where A549 were only significantly affected at 50 µg ml\(^{-1}\). In contrast, the tight junctions were not affected by the NP (Fig. 13(a)) [244].

Moreover, a study employing the interaction between different food matrices and TiO\(_2\) and ZnO NPs (Fig. 13(b)) found that depending on the nanomaterial and the food matrix, it can change the following parameters such as intermolecular interactions and the size of the nanomaterial through the GI tract. These relations could finally modify the toxicological responses induced by NPs after oral exposure [243,245]. It is essential to mention that the doses administered in most studies were excessive compared to what is required in regular human daily consumption. In vitro studies utilizing NPs for oral consumption might lead to uncertain information of toxicity of NPs, and it is necessary to assess the synergistic effects of NPs in a complex system [246].

As mentioned, Zn-associated NPs are in their early life in mineral nutrition, and additional research is required. Further research should be directed to find the optimum levels of Zn NPs that can present better performance and benefits. Evaluation of toxic effects needs a more significant number of studies and the most prolonged period to standardize both the positive and possible adverse effects. In addition, the financial side and stability of the produced mineral NPs beneath normal storage conditions should also be extensively studied. The biological challenges of utilizing metal oxide NPs for efficient antimicrobial activity and scale-up manufacturing for clinical applications are shown in Fig. 13(c) [247]. Henceforth, it is vital to mention that a careful study is required to determine the exact relationship between morphological properties of these nanomaterials and their dissolution, transport, metabolism, oxidative impacts, biotoxicity, and other biological effects. Further experiments should be conducted to characterize the differences between nanosized Zn behavior in animal models and human GI tracts (or different routes of administration). Also, it is imperative to understand the implications of its effect on the gut microbiota and its possible outcome in respiratory virus infection.

5. Immunomodulatory properties of the Zn nanomaterials

Nanotechnology strategies have contributed as advanced solutions for the treatment against emergent viruses like H1N1, MERS, and SARS-CoV-2 [7]. Apart from the attractive applications mentioned earlier in this report, such as diagnosis, disinfection,
Fig. 13. Potential toxicological mechanisms of administration of Zn nanomaterials. (a) Metabolic activity (A), LDH (cytotoxicity) (B), and release and ROS generation (C) after ZnO NP exposure in different lung epithelia cells [244]. Copyright Royal Society of Chemistry. (b) Metal oxide nanoparticles have efficient antimicrobial activity, but many biological challenges have restricted their application and hurdles to scale-up manufacture for the clinic [247]. Copyright John Wiley & Sons 2019. (c) Effect of food on orally ingested nanoparticle behaviors in the simulated digestive tract. Copyright Europe PMC 2020 [245].
and supplementation, ZnO NPs have attractive immunomodulatory properties suggesting their potential use as an adjuvant.

First of all, the immune system is interconnected between innate and adaptative responses. The innate response includes macrophages that engulf pathogens and natural killer (NK) cells that induce cell death in infected host cells. For the adaptative immune system, pathogens or antigens move to immune tissues, like lymph nodes (LN), where antigen-presenting cells (APCs), like dendritic cells (DCs), process and present the antigen on their surface by displaying specialized proteins called major histocompatibility complexes (MHCs) (Fig. 14) [248]. NPs can participate in immunomodulation through the release of various cytokines and growth factors [249]. Specifically, ZnO NPs can induce the expression of immunoregulatory cytokines, which is also a relevant consideration for potential use in biomedical applications, as many current treatments for human disease function by manipulating and controlling components of the immune response [250]. The immunomodulatory response has been associated with systemic deposition of ZnO NPs in the tissues that can cause classic pulmonary oxidative inflammatory responses [251]. For instance, Saptarshi et al. [252] have reported the potential of ZnO NPs to cause pulmonary immunomodulation when exposed via intranasal instillation, highlighting that the cytotoxicity depends on NPs concentration, size, and time of exposition. As mentioned earlier, ZnO NP cytotoxicity has been reported in vitro test systems, including immune cells, lung epithelial cells, etc. [253-256]. The reports suggest that ZnO NP cytotoxicity is mainly associated with ROS generation [256-258]. In a few studies, ZnO NPs are shown to increase the expression of inflammatory markers (ICAM-1, IL-8, and MCP-1) [259] and to enhance the production of cytokines (IFN-γ, TNF, and IL-12) [250].

On the other hand, Poon et al. [260] studied nano- and bulk-sized ZnO NPs, showing their effect on cell metabolisms and gene expression through transcriptomic analyses associated with exposure to the maximum sub-lethal dose to characterize the early immune signaling and material-specific adaptative responses. They found that exposure to ZnO NPs induced the expression of several innate and adaptative immunity pathways.

According to the above discussion, the search for effective adjuvants for improving the immune response is vital (as a form of contention) against emergent respiratory diseases that have not been controlled. Several reports indicate that ZnO NPs have adjuvant properties [249,261-264] and describe the mechanisms involved in adjuvant response induced in cell lines [261]. It is mentioned before, the stimulatory effects are mediated by intracellular Zn2+ dissolution in the lysosomes and ROS generation, resulting in the release of inflammatory cytokines and cellular activation of immune cells [264]. Then, ZnO NPs could participate as either a delivery system to enhance antigen processing or an immune-stimulating adjuvant. For instance, ZnO NPs have also been found to show adjuvant effect via activating the innate immune system and recruiting APCs (CD19+ MHC class-II+ B cells and activated macrophages), lymphocytes, eosinophils, mast cells, and neutrophils [249]. However, these NPs boost the secondary immune response leading to the exacerbation of inflammation characterized by enhanced expression of Cox2, MMP 9, and PGE2 [261].

To explore ZnO NPs intrinsic adjuvant-like properties and immunomodulatory functions, Afroz et al. [262] designed a novel mesoporous ZnO nanocapsule (mZnO) to investigate its immunomodulatory properties by using Ova loaded mZnO-nanocapsules [mZnO(Ova)] in a mice model. Their findings showed that mZnO(Ova) administration steered the enhanced expansion of antigen-specific T-cells and induction of IFN-γ producing effector CD4+ and CD8+ T-cells. Also, antigen-specific IgG levels are enriched in both the serum and lymph nodes of mZnO(Ova) immunized mice. Further, they noticed a substantial increase in serum IgG2a or IgG2b levels and IFN-γ secretion in Ova restimulated splenocytes from mZnO(Ova) immunized mice. Given these features in combination with its immunomodulatory characteristics reinforces the idea that ZnO NPs could be used as an effective antigen-adjuvant platform for developing novel nano-based vaccines against multiple diseases, including respiratory viruses.

Mucosal immunization via the respiratory tract mimics the natural entry of antigens or viruses, such as the case of respiratory viruses like SARS-CoV-2, that usually have the first contact with the nasal or pulmonary mucosa when entering the human body. An earlier study using different spray-dried ZnO formulations to test the aerodynamic performances and immunological response, shown promising properties about mucosal vaccination via the respiratory tract, via increasing the expression of CD80, CD86, and MHC-II, which indicated an activation of the dendritic cells, proposing the use of ZnO for respiratory vaccination [265].
Despite the properties mentioned above, the utilization of Zn NPs or Zn\(^{2+}\) in the vaccine development against virus-like H1N1, H5N1, MERS, or SARS-CoV-2 has not been fully explored even when the nanotechnology has been valuable in the COVID-19 pandemic. Until now, the greatest approach in this sense could be the utilization of Zn in the stabilization of liposomes, taking into account that they are essential components in some of the authorized vaccines (Moderna [266] and Biotech/Pfizer [266]).

Notably, the use of lipid nanoparticles (LNPs) is one of the most promising vaccine platforms, which are highly efficient in encapsulating DNA- or RNA- based immunogens or antibodies[55]. The new generation of nano vaccines coupling LNPs and metal ions, like Zn\(^{2+}\), Mn\(^{2+}\), Co\(^{2+}\), by functionalizing PoP-bilayers or another biomolecule is a promising alternative. For instance, Federizon et al. (2021) reported that Co\(^{2+}\) incorporation into liposomes could increase the immune response, proposing it as a new vaccine

Fig. 15. Zinc participation for stabilizing liposomes as a strategy in the nanovaccines development against SARS-CoV-2. Schematic representation showing the coupling of metallic ions with liposome nanoparticles (LNP) and surface functionalization using SARS-CoV-2 Spike protein (PDB code 6XS6), which may be useful in a novel development of nanovaccines.

Fig. 16. Future perspectives of nano Zn and its compounds for COVID-19 detection and treatment methods.
candidate that warrants further investigation since attaching polypeptides to lipid bilayers is often indirect, ineffective, and can represent a substantial bottleneck in forming functionalized lipid-based materials [267-269]. Consequently, novel prototypes of nanoengineered particles such as the LNPs complex with metallic cations like Zn²⁺, with its intrinsic viricidal activity [157], could be demanding (Fig. 15). More investigation is required to clarify how this novel nanosystem could increase human immune response in the same way as previously observed in animal models [148]. The main limitation of nano vaccines coupling LNPs as a delivery system is their poor stability in gastric juice, high manufacturing cost, and inactivation of phospholipid membrane integrity [270]. However, after coupling with metal ions, their stability can be further improved, and it could open the pathway for their mass production, providing an excellent choice of delivering vaccines. Therefore, nano vaccines research is inherently a field in which more challenges demand our attention. It could help develop more efficient and safer vaccines, mainly directed to people with specific comorbidities.

6. Conclusions and future perspectives

The use of nanomaterials as Zn and its compounds has provided an essential contribution to the management of COVID-19. Some of the significant conclusions from the present study and the future challenges for applying nanostructured Zn and its compounds to combat respiratory viruses are highlighted in Fig. 16. One of the main concerns about COVID-19 is the diagnosis and detection of infected patients. Detection strategies are focused on molecular techniques, which are expensive and time-consuming; however, the need for rapid detection methods is one of the most critical control strategies. The development of nanomaterials as sensor platforms can reduce detection time, reagents, and infrastructure. The use of Zn-based nanostructures to detect the principal biomarkers of the SARS-CoV-2 remains an ongoing challenge, where the development of selective and sensitive devices is the primary goal. The former application of ZnO nanostructures for the effective detection of multiple biomarkers, including viruses, suggests the potential use of these systems for developing electrochemical and optical biosensing platforms for COVID-19 surveillance. However, it is crucial to consider that the main limitations are the costs of fabrication and mass-scale production of these biosensors based on nanotechnology. Additionally, for the successful implementation of biosensors, different analytes must be conserved and transported on distant, isolated zones, which could significantly hinder the implication of these upcoming technologies.

To impede virus propagation, the increasing demand for self-sterilized coating requires alternative materials to fulfill this issue. In this respect, the use of Zn/ZnO nanostructures has exhibited significant antiviral activity against several viruses such as influenza and coronaviruses. As illustrated in Fig. 16, the cytotoxic study of this material, along with the potential scalability, is the leftover task. On the other hand, the role of Zn as a supplement has shown benefits such as modulation of the immune response and antiviral activity against SARS-CoV-2 and other respiratory viruses, although more studies are still needed. Therefore, it is necessary to determine the relationship between nano Zn materials and their biotoxicity, dissolution, transport, metabolism, oxidative impacts, and other biological effects in the human body. Until now, ZnO NPs have not been used to develop vaccines; however, their potential adjuvant properties have been explored, producing several innate and adaptive immunity responses. Taking this into consideration, the addition of metal ions to stabilized liposomes could be a way to develop novel platforms that may include Zn or ZnO NPs, since the global research efforts develop on the convergence of specific vaccines. This could help develop more efficient and safer vaccines coupling LNP and metal ions, even though its final effect on the immune response is still under study. It is noteworthy to mention that this materials appealing properties can be extrapolated to new clinical challenges and can also be used in emergency health situations like the COVID-19 pandemic. Overall, the implementation of Zn-based nanomaterials could impact technological development, thus contributing to well-being and economic growth.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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References

[1] C.D. (COVID-19), WHO, (2021). https://covid19.who.int/?gclid= CJwvKAjsw1X0GRBBVkeAwshF7Bw8jy00_BuVzvLxsvbJuzzhu90TH_Occ4s5gUeuUp YRyEpEjJl03ZFRJYxoC0cM0QAVD_Bw.
[2] S. Jain, L. Finelli, M.W. Shaw, S. Lindstrom, R.J. Garten, L.V. Guavera, X. Xu, C. B. Bridges, T.M. Uyeki, Emergence of a Novel Swine-Origin Influenza A (H1N1) Virus in Humans, N. Engl. J. Med. 360 (2009) 2605–2615, https://doi.org/10.1056/nejmoa0993810.
[3] J.W. LeDuc, M.A. Barry, SARS, The First Pandemic of the 21st Century, Emerg. Infect. Dis. 10 (2004) e26. 10.3201/eid1011.040797_02.
[4] S. Talebian, G.G. Wallace, A. Schroeder, F. Stellacci, J. Conde, Nanotechnology—overview and perspectives, J. Control. Release. 240 (2016) 24–37, https://doi.org/10.1016/j.jconrel.2015.10.012.
[5] W.C. Huang, S. Zhou, X. He, K. Chiem, M.T. Mabrouk, R.H. Nissly, L.M. Bird, M. Strauss, S. Sambharia, J. Ortega, E.A. Wohlfert, L. Martinez-Sobrado, S.V. Kuchipudi, B.A. Davidson, J.F. Lovell, SARS-CoV-2 RRBD neutralizing antibody induction is enhanced by particulate vaccination, Adv. Mater. 2005637 (2020) 1–11, https://doi.org/10.1002/adma.202005637.
[6] T. Hu, Y. Liu, M. Zhao, Q. Zhuang, L. Xu, Q. He, A comparison of COVID-19, SARS and MERS, PeerJ 8 (2020) 1–30, https://doi.org/10.7717/peerj.9725.
[7] S. Dawre, S. Maru, Human respiratory viral infections: current status and future prospects of nanotechnology-based approaches for prophylaxis and treatment, Life Sci. 278 (2021), https://doi.org/10.1016/j.lfs.2021.119561.119561.
[8] M. Yu, J. Wu, J. Shi, O.C. Farokhzad, Nanotechnology for protein delivery: overview and perspectives, J. Control. Release. 240 (2016) 24–37, https://doi.org/10.1016/j.jconrel.2015.10.012.
[9] W.C. Huang, S. Zhou, X. He, K. Chiem, M.T. Mabrouk, R.H. Nissly, L.M. Bird, M. Strauss, S. Sambharia, J. Ortega, E.A. Wohlfert, L. Martinez-Sobrado, S.V. Kuchipudi, B.A. Davidson, J.F. Lovell, SARS-CoV-2 RRBD neutralizing antibody induction is enhanced by particulate vaccination, Adv. Mater. 2005637 (2020) 1–11, https://doi.org/10.1002/adma.202005637.
[10] M.W. Li, Y. Yi, X. Luo, N. Xiong, Y. Liu, S. Li, Q. Sun, Y. Wang, B. Hu, W. Chen, Y. Zhang, J. Wang, B. Huang, Y. Lin, J. Yang, W. Cai, X. Wang, J. Cheng, Z. Chen, K. Sun, W. Pan, Z. Zhan, L. Chen, F. Ye, Development and clinical application of a
rapid IgM-IgG combined antibody test for SARS-CoV-2 infection diagnosis, J. Med. Virol. 92 (2020) 1518–1524, https://doi.org/10.1002/jmv.25727.

[14] H. Zhao, F. Liu, W. Xie, T.C. Zhou, J. OuYang, L. Jin, H. Li, C.V. Zhao, L. Zhang, J. Wei, Y.P. Zhang, C. Liu, Ultrasensitive sandwich-type electrochemical sensor for SARS-CoV-2 from the infected COVID-19 patients using a smartphone, Sens. Actuators, B Chem. 327 (2021), https://doi.org/10.1016/j.snb.2021.129066.

[15] E.V.R. Campos, A.E.S. Pereira, J.L. De Oliveira, L.B. Carvalho, M. Guigler-Casagrande, R. De Lima, F.F. Racetto, How can nanotechnology help to combat COVID-19? Opportunities and urgent need, J. Nanobiotechnol. 18 (2020) 1–23, https://doi.org/10.1186/s12985-020-00363-9.

[16] A.M. Shaw, C. Hyde, B. Merrick, P. James-Pemberton, B.K. Squires, R.V. Olkhov, A.M. Shaw, C. Hyde, B. Merrick, P. James-Pemberton, B.K. Squires, R.V. Olkhov, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[17] A. Karami, M. Hasani, F. Azizi Jalilian, R. Ezati, Conventional PCR assisted enhanced plasmonic detection scheme of SARS-CoV-2 Spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[18] A.M. Shaw, C. Hyde, B. Merrick, P. James-Pemberton, B.K. Squires, R.V. Olkhov, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[19] D. Zhang, X. Zhang, R. Ma, X. Wang, X. Zhang, X. Huang, Y. Liu, G. Li, J. Qi, Y. Zhu, J. Li, Ultra-fast and onsite interrogation of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) in waters via surface enhanced Raman scattering (SERS), Water Res. 200 (2021), https://doi.org/10.1016/j.watres.2021.117243.117243.

[20] C. Wang, X. Yang, B. Gu, C. Hu, L. Liu, Z. Zhou, L. Shi, X. Cheng, S. Wang, C. Wang, B. Gu, S. Wang, Sensitive and simultaneous detection of SARS-CoV-2-Specific IgM/IgG using lateral flow immunoassay based on dual-ratio quantum dot nanobeads, Anal. Chem. 92 (2020) 15542–15549, https://doi.org/10.1021/acs.analchem.0c04384.

[21] Z. Zhao, H. Cui, W. Wang, X. Xu, W. Zhou, X. Xu, A simple magnetic nanoparticles-based viral RNA extraction method for efficient detection of SARS-CoV-2, 518055 (2020). 10.1101/2020.02.22.961268.

[22] A. Karami, M. Hasani, F. Azizi Jalilian, R. Ezati, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[23] A. Karami, M. Hasani, F. Azizi Jalilian, R. Ezati, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[24] A. Karami, M. Hasani, F. Azizi Jalilian, R. Ezati, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.

[25] A. Karami, M. Hasani, F. Azizi Jalilian, R. Ezati, Conventional PCR assisted enhanced plasmonic detection scheme of COVID-19 SARS-CoV-2 spike protein, Adv. Theory Simul. 3 (2020) 1–8, https://doi.org/10.1002/axtt.2019.01.047.
[132] Y. Xie, Y. He, P.L. Irwin, T. Jin, X. Shi, Antiviral activity and mechanism of action of zinc oxide nanoparticles against Campylobacter jejuni, Appl. Environ. Microbiol. 77 (2011) 2325–2331, https://doi.org/10.1128/AEM.01449-10.

[133] S. Sharmin, M.M. Rahman, C. Sarkar, O. Atofani, M.T. Islam, O.S. Adeyemi, Nanoparticles as antimicrobial and antiviral agents: a literature-based perspective study, Helioven. 7 (2021), https://doi.org/10.1016/j.heliov.2021.01.001.

[134] S.H. Cha, J. Hong, M. McGuffie, B. Yeom, J.S. Vanpepe, N.A. Kotov, Shape-dependent biomimetic inhibition of enzyme by nanoparticles and their antimicrobial activity, ACS Nano 9 (2015) 9907–9915, https://doi.org/10.1021/acsnano.5b04856.

[135] C.L. de Dicostato, C.P. Vidal, I. Falcón, S. Páez, M. Matsuura, L. Yanagida, C. Alves, Y.-H. Pan, C.-L. Chiu, H.-C. Kao, Cholesterol-mediated cytotoxicity and inhibition of SARS-CoV-2 infection by ZnO nanoparticles: green synthesis, characterization, and antimicrobial activity, Nanomedicine. 13 (2017) 1861–1871, https://doi.org/10.2217/nnm-2017-0089.

[136] A. Adhikari, U. Pal, S. Rayan, S. Mondal, R. Ghosh, S. Darbar, T. Saha-Dasgupta, S.K. Ray, S.K. Pal, Nanocatalytic firesman prevents COVID-19 spread through expelling respiratory droplets: a combined computational, spectroscopic, and antimicrobial study, ACS Appl. Mater. Interfaces 4 (2021) 5471–5484, https://doi.org/10.1021/acsami.1c03228.

[137] S.M. El-Megharbel, A. Alsawat, F.A. Al-Salmi, R.Z. Hamza, Utilizing of (Zinc oxide) nanoparticles—a nano weapon to fight against herpes simplex virus-1, Antiviral Res. 92 (2011) 305–312, https://doi.org/10.1016/j.antiviral.2011.08.017.

[138] M. Kar, H.A. Khan, A. Panwar, S.S. Bais, S. Basak, R. Goel, S. Sopory, G.R. Medegishi, Zinc chelation specifically inhibits early stages of dengue virus replication by activation of NF-kappa B and induction of antiviral response in HepG2 cells, Front. Immunol. 10 (2019) 2347, https://doi.org/10.3389/fimmu.2019.02347.
C. Gutiérrez Rodelo, R.A. Salinas, E. Armenta Jaime et al. Coordination Chemistry Reviews 457 (2022) 214402

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D. Jothimani, E. Kailasam, S. Danielraj, B. Nallathambi, H. Ramachandran, P. M. Jarosz, M. Olbert, G. Wyszogrodzka, K. Młyniec, T. Librowski, Antioxidant N.Z. Gammoh, L. Rink, Zinc in infection and inflammation, Nutrients 9 (2017)

166

S. Mehta, Nutritional status and COVID-19: an opportunity for lasting 2021 N.I. of H. National Institutes of Health, 2021, COVID-19 Treatment Scott A Read, Peran Seng dalam Kekebalan Antiviral, Univ. Sydney Westmead 2019, 145

R. Wegmüller, F. Tay, C. Zeder, M. Brnic’, R.F. Hurrell, Zinc absorption by young 2021, PLoS ONE. 16 (2021) 1–12.

S. Rerksuppaphol, L. Rerksuppaphol, A randomized controlled trial of 2036, 369. 10.1016/j.sj.2002.09.027.

D.K. Perry, M.J. Smyth, H.R. Stennicke, G.S. Salvesen, P. Duriez, G.G. Poirier, A. Sabin, Zinc is a zinc-dependent inhibitor of the 3C-like serine-protease, an early zinc transporter candidate for zinc as a putative target for zinc in the inhibition of apoptosis, J. Biol. Chem. 272 (1997)

I. Wittes, B. Rolles, L. Rink, The zinc deficiency and zinc supplementation in young adults from supplement zinc citrate is compatible with that from zinc gluconate and higher than from zinc oxide, J. Nutr. 144 (2014) 132–136, https://doi.org/10.3944/jn.113.918417.

B. Rolles, M. Maywald, L. Rink, Influence of zinc deficiency and supplementation on NK cell cytotoxicity, J. Pestic. Sci. 48 (2013) 322–328, https://doi.org/10.1002/jps.23707.

E. Rosenkranz, C.H.D. Metz, M. Maywald, R.D. Hilgers, I. Wittes, S. Senff, H. Haase, M. Jager, M. Ott, R. Aspinall, B. Plumakers, L. Rink, Zinc deficiency reactivates inducible T cells by inhibition of Sirt1-deacetylase in mixed lymphocyte cultures, Mol. Nutr. Food Res. 60 (2016) 656–669, https://doi.org/10.1007/s00424-016-1917-3.

A. Pernalek, A novel form of host defence: membrane protection by Ca2+ dependent RNA-polymerase and 3C-like proteinase enzymes, Int. J. Mol. Med. 28 2009, 2985S-2989S. 10.1093/jn/133.9.2985s.

D.K. Perry, M.J. Smyth, H.R. Stennicke, G.S. Salvesen, P. Duriez, G.G. Poirier, A. Sabin, Zinc is a zinc-dependent inhibitor of the 3C-like serine-protease, an early zinc transporter candidate for zinc as a putative target for zinc in the inhibition of apoptosis, J. Biol. Chem. 272 (1997)

I. Wittes, B. Rolles, L. Rink, The potential impact of zinc supplementation on COVID-19 pathogenesis, Front. Immunol. 11 (2020) 1712, https://doi.org/10.3389/fimmu.2020.01712.

S. Reed, H. Skak, Dietary zinc alters the microbiota and decreases resistance to Clostridium difficile infection, Nat. Med. 22 (2016) 1330–1334, https://doi.org/10.1038/nm.4174.

S. Meiia, C. Sammy, M. Suckovskich, R. Ghun, O. Koen, E. Tako, Chronic zinc deficiency alters chick gut microbiota composition and function, Nutrients 7 (2015)

O. O’Callaghan, The effects of coronavirus on human nasal ciliated respiratory 2020, 624, https://doi.org/10.3365/e090562.

D. Dhar, A. Mohanty, Gut microbiota and Covid-19- possible link and implications, Viruses 2 (2020) 85–285, https://doi.org/10.3390/v2018018108.

S.J.M. Osendarp, H. Prabhakar, G.J. Fuchs, J.M.A. van Raaij, M. Mahfud, F. Mol, T. Mostad, S. Black, Immunization with the heptavalent pneumococcal conjugate vaccine protects Bangladeshi infants and effects of zinc supplementation, Vaccine 25 (2007) 3347–3354, https://doi.org/10.1016/j.vaccine.2007.01.001.

D. Zhao, S. Yu, G. Sun, H. xin Zhang, Y. Chen, M. Guo, Zinc Deficiency Promoted Fibrosis via ROS and TIMP/MMPs in the Myocardium of Mice, Biol. Trace Elem. Res. 196 (2020) 145–190, https://doi.org/10.1007/s12011-019-01902-4.

M.A. Chilvers, M. McKean, A. Rutman, B.S. Myint, M. Silverman, C. O’Callaghan, The effects of coronavirus on human nasal ciliated respiratory epithelium, Eur. Respir. J. 18 (2001) 965–987, https://doi.org/10.1183/09031901.0009301.

M. Singh, R.R. Das, Zinc for the common cold, Cochrane Database Syst. Rev. 2013 (2013), https://doi.org/10.1002/14651858.CD001364.pub4.
[206] M.C. Sportelli, M. Izzi, E.A. Kukushkina, S.I. Hossain, R.A. Picca, N. Ditaranto, N. I. Abdulhamid, F.W.J. Beck, S. Millard, X. Chen, A. Prasad, Effect of zinc inhibitory effects of epigallocatechin gallate (EGCG) combined with zinc. Coordination Chemistry Reviews 457 (2022) 214402.

[207] K.S. Siddiqi, A. ur Rahman, Tajuddin, A. Husen, Properties of zinc oxide nanoparticles for the prevention of acute respiratory infections in infants: a randomized, double-blind, placebo-controlled clinical trial. JAMA Network, American Medical Association, (2021).

[208] D. Hulisz, Efficacy of zinc against common cold viruses: an overview, J. Am. Nutr. 32 (2013) 193–199, https://doi.org/10.1016/j.clnu.2012.08.018.

[209] D. Mahalanabis, M. Lahiri, D. Paul, S. Gupta, A. Gupta, M.A. Wahed, M.A. Khaled, Randomized, double-blind, placebo-controlled clinical trial of the efficacy of treatment with zinc or vitamin A in infants and young children with severe acute lower respiratory infection, Am. J. Clin. Nutr. 79 (2004) 430–436, https://doi.org/10.1093/ajcn/79.4.430.

[210] U.H. Shah, A.K. Abu-Shaheen, M.A. Malik, S. Alam, M. Riaz, M.A. Al-Tannir, The toxicology of zinc chloride smoke producing bombs and screens. Biochem. 45 (2021),https://doi.org/10.1111/jfbc.13604.

[211] S. Thomas, D. Patel, B. Bittel, K. Wolski, Q. Wang, A. Kumar, Z.J. Il’Giovine, R. Mehra, C. McWilliams, S.E. Nissen, M.Y. Desai, Effect of high-dose zinc and ascorbic acid supplementation vs usual care on symptom length and reduction among ambulatory patients with SARS-CoV-2 infection: the COVID A to Z randomized clinical trial. JAMA Netw. Open. 4 (2021) e210369–e210369. 10.1001/jamanetworkopen.2021.0369.

[212] M.P. Vinardell, H. Llanas, W. J, B. A, T. R, T. R, H. H, G. GP, D. M, W. A, D. RK, Two-dimensional antiviral zinc oxide nanoparticles mediated by hesperidin and in silico simulations from sunscreens containing zinc oxide nanoparticles: a review. Nanotoxicology 10 (2016) 720–727. 10.3109/17435390.2015.1113322.

[213] S.W. Mcpherson, J.E. Keunen, A.C. Bird, E.Y. Chew, F.J. Van Kuijk, EDITORIAL investigation oral zinc as a prophylactic treatment for those at risk for COVID-19, (2020). 10.1016/j.jfs.2020.04.038.

[214] L. Wang, J. Liu, M. N. Kayow, L.I. Hird, H. Zhang, Ameliorative effect of Alloba chiensis synthesized ZnO-NPs on Mycoplasma pneumoniae infected pneumonia mice model, Microb. Pathiol. 141 (2020), https://doi.org/10.1016/J.MICPATH.2019.103960.

[215] D. Hulisz, E. Simeoni, V. Angelii, G.J. Randolph, C.P. O’Neil, L. Lee, M.A. Swartz, J.A. Hubbell, Exploiting lympathic transport and complement activation in nanoparticle vaccines, Nat. Biotechnol. 25 (2007) 1277–1285, https://doi.org/10.1038/nbt1332.

[216] S.W. Mcpherson, J.E. Keunen, A.C. Bird, E.Y. Chew, F.J. Van Kuijk, EDITORIAL investigation oral zinc as a prophylactic treatment for those at risk for COVID-19, (2020). 10.1016/j.jfs.2020.04.038.

[217] S.T. Reddy, A.J. Van Der Vlies, E. Simeoni, V. Angelii, G.J. Randolph, C.P. O’Neil, L. Lee, M.A. Swartz, J.A. Hubbell, Exploiting lympathic transport and complement activation in nanoparticle vaccines, Nat. Biotechnol. 25 (2007) 1277–1285, https://doi.org/10.1038/nbt1332.

[218] S. Abd-Elsalam, S. Soliman, E.S. Esmail, M. Khalaf, E.F. Mostafa, M.A. Medhat, A. Khaled, A. Abou-Tarek, M.A. Elbayoumi, A. Elshemay, A. Zaki, I.K. Seif, M.Y. Renawy, M. Amin, K. Amer, M.A. El Demelawy, Nanoparticles of ZnO/Beirubene complex contract COVID-19 and respiratory co-bacterial infection in addition to elimination of hydroxylcorticoid toxicity, J. Pharm. Invest. (2021) 1, https://doi.org/10.5408/s10504-021-00544-W.

[219] M.J. Saadh, M.M. Agagg, A. Alboghdally, M. Khashirid, S.M. Aldalmaen, A. Abdelrazek, Silver nanoparticles with epigallocatechin gallate and zinc sulphate significantly inhibits avian influenza A virus H9N2. Microb. Pathiol. 158 (2021), https://doi.org/10.1016/j.micpath.2021.105071.

[220] S. Syama, P.J. Sreekanth, H.K. Varma, P.V. Mohanan, Zinc oxide nanoparticles induced oxidative stress in mouse bone marrow mesenchymal stem cells, Toxicol. Mech. Methods. 24 (2014) 644–653, https://doi.org/10.3109/153735613X1392296.

[221] S. Syama, P.J. Sreekanth, H.K. Varma, P.V. Mohanan, Zinc oxide nanoparticles induced oxidative stress in mouse bone marrow mesenchymal stem cells, Toxicol. Mech. Methods. 24 (2014) 644–653, https://doi.org/10.3109/153735613X1392296.

[222] S. Abd-Elsalam, S. Soliman, E.S. Esmail, M. Khalaf, E.F. Mostafa, M.A. Medhat, A. Khaled, A. Abou-Tarek, M.A. Elbayoumi, A. Elshemay, A. Zaki, I.K. Seif, M.Y. Renawy, M. Amin, K. Amer, M.A. El Demelawy, Nanoparticles of ZnO/Beirubene complex contract COVID-19 and respiratory co-bacterial infection in addition to elimination of hydroxylcorticoid toxicity, J. Pharm. Invest. (2021) 1, https://doi.org/10.5408/s10504-021-00544-W.

[223] M.J. Saadh, M.M. Agagg, A. Alboghdally, M. Khashirid, S.M. Aldalmaen, A. Abdelrazek, Silver nanoparticles with epigallocatechin gallate and zinc sulphate significantly inhibits avian influenza A virus H9N2. Microb. Pathiol. 158 (2021), https://doi.org/10.1016/j.micpath.2021.105071.
[238] C. McCracken, P.K. Dutta, W.J. Waldman, Critical assessment of toxicological effects of ingested nanoparticles, Environ. Sci. Nano 3 (2016) 256–282, https://doi.org/10.1039/c6nj00242e.

[239] V. Tattineni, J.Y. An, M.R. Leffew, J.Y. An, M.R. Leffew, S.A. Mahesh, Anemia from A to zinc: oxidative stress and genotoxicity induced by four typical nanomaterials: the role of particle size, shape and composition, J. Appl. Toxicol. 29 (2009) 69–78, https://doi.org/10.1002/jat.1385.

[240] A. Gojova, B. Guo, R.S. Kota, J.C. Rutledge, I.M. Kennedy, A.I. Barakat, Induction of inflammation in vascular endothelial cells by metal oxide nanoparticles: effect of particle composition, Environ. Health Perspect. 115 (2007) 403–409, https://doi.org/10.1289/ehp.8497.

[241] W.L. Poon, H. Alenius, J. Ndika, V. Fortino, V. Kolhinen, A. Mešcéríakovas, M. Wang, D. Greco, A. Lähde, J. Kokkinemi, J.C.Y. Lee, H. El-Nezami, P. Karisola, Nano-sized zinc oxide and silver, but not titanium dioxide, induce innate and adaptive immunity and antiviral response in differentiated THP-1 cells, Nanotoxicology 11 (2017) 936–951, https://doi.org/10.1080/17435390.2017.1382600.

[242] R. Roy, D. Kumar, A. Sharma, P. Gupta, B.P. Chaudhari, A. Tripathi, M. Das, P.D. Dwivedi, ZnO nanoparticles induced adjuvant effect via toll-like receptors and Snr signaling in Balb/c mice, Toxicol. Lett. 230 (2014) 421–433, https://doi.org/10.1016/j.toxlet.2014.08.008.

[243] S. Afroz, H. Medhi, S. Mithin, S. Batto, J. Giddaluru, K. Kumar, P. Paik, N. Khan, Mesoporous ZnO nanocapsules for the induction of enhanced antigen-specific immunological responses, Nanoscale. 9 (2017) 14641–14653, https://doi.org/10.1039/c7nr03697c.

[244] R. Roy, S. Kumar, A.K. Verma, A. Sharma, B.P. Chaudhari, A. Tripathi, M. Das, P. Dwivedi, Zinc oxide nanoparticles provide an adjuvant effect to oseltamivir via a th2 response in Balb/c mice, Int. Immunol. 26 (2014) 159–172, https://doi.org/10.1093/intimm/dxu053.

[245] P. Sharma, N.Y. Jang, J.W. Lee, B.C. Park, Y.K. Kim, N.H. Cho, Application of ZnO-based nanocomposites for vaccines and cancer immunotherapy, Pharmaceuticals 11 (2019) 6–10, https://doi.org/10.3390/pharmaceutics11040493.

[246] M.G. Hellfrisch, ZINC OXIDE IN FORMULATION, 2021.

[247] L.A. Jackson, E.J. Anderson, N.G. Rouphael, M. Makhene, R.N. Buchanan, R. Pikaart-Tautges, J.E. Ledgerwood, B.S. Graham, J.H. Beigel, An mRNA Vaccine against SARS-CoV-2 — Preliminary Report, N. Engl. J. Med. 383 (2020) 1920–1931, https://doi.org/10.1056/nejmoa2022483.

[248] W.C. Huang, B. Deng, A. Selfoff, J. Ortega, C.A. Long, R.V. Suresh, J.F. Lovell, Antibody response of a particle-inducing, liposome vaccine admixed with a Pfs230 fragment, NPJ Vaccines 5 (2020) 1–9, https://doi.org/10.1038/s41541-020-0173-x.

[249] J. Fedorcz, C.G.T. Feugmo, H.W. Huang, X. He, K. Miura, A. Razi, J. Ortega, M. Karttunen, J.F. Lovell, Experimental and computational observations of immunogenic bovine porphyrin lipid bilayers: nanodomain-enhanced antigen association, Pharmaceutics 13 (2021) 1–17, https://doi.org/10.3390/pharmaceutics13010008.

[250] S. Shao, J. Geng, H. Yi, S. Gogia, S. Neelameghan, A. Jacobs, J. Lovell, Functionalization of cobalt porphyrin-phospholipid bilayers with His-tagged ligands and antigens, Nat Chem. 7 (2015) 438–446, https://doi.org/10.1038/nchem.2236.

[251] K. Khorasvi-Darani, M.R. Mozaafari, Nanoparticle Potential in Nanotherapy: A Concise Overview, n.d.

[252] S. Mahari, A. Roberts, D. Shahdeo, S. Gandhi, ecoCovSens-ultrasensitive novel in-house built printed circuit board based electrochemical device for rapid detection of nCoV-19 antigen, a spike protein domain 1 of SARS-CoV-2, BioRxiv (2020), https://doi.org/10.1101/2020.04.24.059204.

[253] M. Xiao, L. Huang, X. Dong, K. Xie, H. Shen, C. Huang, W. Xiao, M. Jin, Y. Tang, Integration of a 3D-printed read-out platform with a quantum dot-based immunosensor for detection of the avian influenza A (H7N9) virus, Analyst 144 (2019) 2594–2603, https://doi.org/10.1039/c9an02336a.

[254] S.Y. Chang, K.Y. Huang, T.L. Chao, H.C. Kao, Y.H. Pang, L. Lu, C.L. Chiu, H.C. Huang, T.J.R. Cheng, J.M. Fang, P.C. Yang, Nanoparticle composite TPNT1 is effective against SARS-CoV-2 and influenza viruses, Sci. Rep. 11 (2021) 1–13, https://doi.org/10.1038/s41598-021-87254-3.