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A comprehensive review on current COVID-19 detection methods: From lab care to point of care diagnosis

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ABSTRACT

Without a doubt, the current global pandemic affects all walks of our life. It affected almost every age group all over the world with a disease named COVID-19, declared as a global pandemic by WHO in early 2020. Due to the high transmission and moderate mortality rate of this virus, it is also regarded as the panic-zone virus. This potentially deadly virus has pointed out the significance of COVID-19 research. Due to the rapid transmission of COVID-19, early detection is very crucial. Presently, there are different conventional techniques available for coronavirus detection like CT-scan, PCR, Sequencing, CRISPR, ELISA, LFA, LAMP. The urgent need for rapid, accurate, and cost-effective detection and the requirement to cut off shortcomings of traditional detection methods, make scientists realize to advance new technologies. Biosensors are one of the reliable platforms for accurate, early diagnosis. In this article, we have pointed recent diagnosis approaches for COVID-19. The review includes basic virology of SARS-CoV-2 mainly clinical and pathological features. We have also briefly discussed different types of biosensors, their working principles, and current advancement for COVID-19 detection and prevention.

1. Introduction

The novel form of coronavirus, whose existence was recognized in 2019 is commonly known as COVID-19 infection, a highly transmitted disease otherwise called severe acute respiratory syndrome infected every age group throughout the globe [1,2]. The outbreak of this Virus was firstly identified in the Wuhan district (China) from the patient with unusual pneumonia-like symptoms, which later spread uncontrollably throughout the planet [2]. It was identified as COVID-19 (earlier known as 2019-nCov) (novel coronavirus) on January 12, 2020 by WHO (World Health Organization) [2]. ICTV (International Committee on Taxonomy of Viruses) annotated the COVID-19 causing virus as SARS-CoV-2 on Feb 11, 2020 [1]. In terms of transmissibility and outbreak region, SARS-CoV-2 simply transcended previous coronavirus outbreaks with SARS (severe acute respiratory syndrome) and MERS (Middle East respiratory symptoms) [3]. Due to the rate of transmission and an increasing number of cases WHO announced a global pandemic for COVID-19 on March 12, 2020. [4,5]. According to WHO published data 160,813,869 confirmed cases and 3,339,002 deaths were reported till May 15, 2021 [4,5].

Based on its infection rate and impact on the socio-economic condition, the utmost concern is to develop tools, techniques, and strategies to overcome this pandemic [144]. Multiple surveillance and screening tools are still needed to develop for early detection of the novel coronavirus and its mutated versions. In only two-decade five pandemics have already hit our community which is SARS, MERS, Swine Flu, Ebola, and SARS-CoV 2. For that reason, early identification of such viruses will not only prevent the spreading of infection but also helps in the socio-economic benefit of humankind [144]. In recent times during the COVID-19 Pandemic, there is some preexisting diagnostic approach for detection and screening of the virus which is also considered the gold standard in the field of medical microbiology. Previously X-rays were
used to take images of the chest for identification of chest infection during the initial stage of the pandemic when there was a lack of information about novel coronavirus [6]. When the data became available related to structural and genomics study with the help of microscope and sequencing, various molecular tools like PCR (Polymerase chain reaction), serologic tests were developed and optimized for COVID-19 screening. In case of PCR (Polymerase chain reaction) various sequence of the virus was targeted based upon the genomics sequence available in the database with specific primers and probes [7]. In the case of serological test kits, antigen or antibodies from the blood samples were used for diagnosis of the virus [8]. In most of the case, the serological kits available in the market are based on LFA (Lateral flow assay) or ELISA (Enzyme-linked immunosorbent assay) as shown Table 3.

To address the limitations associated with traditional detection methods, multiple physical diagnostic tools have developed in recent times which are mostly based on biosensors due to their beneficial properties like high sensitivity, high specificity, low cost, and easy to use [9]. Biosensors are the device having both organic and inorganic components which analyze and provide qualitative and quantitative information about the specific type of analyte in the provided samples [5]. The organic component of the biosensors are biological elements which can be antibody or aptamer, and the inorganic components are transducers which generates signals in the sensor [10]. Based on these types of components used in biosensors they can be immune-sensors, Enzyme based sensors, Electrochemical sensors, Optical sensors, FRET-based sensors, Colorimetric sensors, Piezoelectric sensors, Magnetoelectric sensors, and others [11]. Advancement in the field of nanotechnology and microfluidics is creating numerous pathways in developing biosensors with multidimensional properties [12,13]. This not only helps in miniaturization in the size of biosensors but also increases the accessibility to common people in the pandemic situation. These physicochemical-based biosensors in the field of diagnostics, aside from been considered as the first line of defense and also in high demand, due to their flexible properties.

These biosensors are already present in the market as rapid diagnostic kits and some are still in the phase of development which are mostly based on the principle of CRISPR technology, microarray, microfluidics, isothermal amplification, and others which are assumed to be promising technologies that can diagnose the virus in the early stage of the infection [14].

This review presents a broad insight into the pathogenic mechanism of SARS-CoV-2 and significant virulence factors. Besides, potential interventions and conventional diagnosis methods to arrest COVID-19 transmission are also discussed. We further pointed up the advancement of recent technologies in the field of biosensors that have sped up the rapid, accurate diagnosis into practice.

2. Details on COVID-19

2.1. COVID-19 and its clinical significance

It is found that several clinical signs are associated with COVID-19 infection wherein most of the case symptoms appear within 3–7 days [15]. Fever, diarrhea, runny nose, pharyngalgia, myalgia-like symptoms appear in the first week which can create hypoxia and dyspnea in the later week of infection. In the critical condition of this infection, the disease can be progressed as ARDS (acute respiratory distress syndrome), septic shock, and coagulation disorder [15]. In some cases, gastrointestinal discomfort, vomiting, depression, shortness of breath is seen [16]. The infection is transmitted due to inhalation of the droplet which can be arisen by sneezing, coughing, or talking with infected Pearson [17]. It can be also spread by direct or indirect contact where the virus can enter inside the body through the mouth, eyes, nose, or other mucosal layers [17]. It is found that viral load is higher in the nasal cavity compared to the throat [3].

Apart from symptomatic, patients can be classified into asymptomatic and presymptomatic. In true asymptomatic cases, COVID-19 positive patients do not experience any symptoms though, in the case of presymptomatic cases, patients may not develop any symptoms before diagnosis but symptoms can come up within 14 days of diagnosing positive [18].

2.2. Basic biology of SARS-CoV-2

Coronaviruses are a class of ss (Single-stranded) RNA containing enveloped viruses as shown in Fig. 1, that possess an evolutionary rate of 4–10 nucleotides per site per year [17]. Corona has been categorized into four broad genera which are the α-coronavirus, β-coronavirus, γ-coronavirus, and δ-coronavirus [3]. The most common example of α-coronavirus is human-associated coronavirus NL63, 229E, Porcine epidemic diarrhea virus [17]. While, beta coronavirus 1, human coronavirus, SARS, SARS-CoV 2 are in beta genera and porcine coronavirus HKU1 is in delta genera [17]. Amidst, α and β-coronaviruses mainly transmit the disease to mammals, γ-coronaviruses affect birds and δ-coronaviruses can transmit to both mammals and birds [3].

SARS-CoV-2 is the causing virus for COVID-19 which is more dangerous than the Middle East respiratory syndrome coronaviruses (MERS-CoV) and SARS-CoV, in terms of pathogenicity [19]. Understanding the virulence and pathogenic mechanism associated with SARS-CoV-2 is the urgent research priority to accelerate COVID-19 therapeutic interventions.

The size of the COVID-19 virus is in the range of 60–140 nm diameter. The genetic material of this virus is comprised of single-stranded positive-sense RNA with almost 29,884 nucleotide sizes [21]. The genome of the virus codes for twenty-seven types of proteins. Some of them are helicase, papain-like protease, 3CL protease, RNA-dependent RNA polymerase, and four structural proteins required during virus assembly, etc. [21]. Almost two-thirds of the SARS-CoV-2 genome consists of ORF1a (13,468 kb) and ORF1b (21,563 kb). These 5′ ended genes express polyprotein 1a and polyprotein 1b. Whilst at the 3′ end SARS-CoV-2 encode structural proteins comprise of spike surface glycoprotein (S), Small envelope protein(E), matrix protein (M), and nucleocapsid protein(N) [19]. Additionally, the SARS-CoV-2 genome also codes for several accessories’ proteins like 3a, 3b, 6, 7a, 7b, 8, 9b, 9c, and 10 as shown in Figs. 1 and 3. These proteins are required during viral replication [20].

2.3. Binding and pathogenicity

Coronaviruses follow a complex receptor-mediated host cell machinery to invade the host cell [17]. For instance, alpha coronavirus human coronavirus NL63 (HCoV-NL63) recognize aminopeptidase N, beta coronavirus like MERS-CoV recognize serine peptidase or dipeptidyl transferase 4 (DPP4) [3]. Mouse hepatitis virus (MHV) mainly interacts with host cell adhesion glycoprotein cell-adhesion glycoprotein carcinoembryonic antigen-related cell adhesion molecule 1 [22,23]. Further, HCoV-NL63, SARS-CoV-2 can recognize angiotensin-converting enzyme 2 (ACE2) in host cells [24,42]. Coronaviruses utilize diverse host receptor recognizing groups for invasion. Other than receptor recognizing coronavirus spike protein machinery carry out multiple viral physiological functions. The particular configuration of spike protein assists the interaction between receptor binding motif (RBM) of S1 protein and ACE2 ectodomain [19].

ACE2 receptors are frequently found in type II alveoli cells of lungs and other tissues. The binding affinity of this receptor SARS-CoV-2 virus spike protein is very high [25]. So, it is considered that the cells which express the ACE2 receptor are the main target for the viral infection as shown in Fig. 2. This ACE2 also expresses cells in oral tissues which suggest the oral cavity can be the route of entry of the virus [26]. Basigin also is known as CD147 encoded by the BSG gene, can be also used by SARS-CoV-2 as an entry site where spike protein binds with this receptor [26]. The attachment of SARS-CoV-2 S protein with host ACE2 activates
cellular TMPRSS2 protease which opens the spike protein and exposes fusion protein and, in that way, the virus is endocytosed inside the host cell [26]. Inside the human cell, the virus uncoats the N protein to release the single-stranded (ss) RNA [27]. Through the activity of replicase and transcriptase gene, viral genetic material replicates that translated later to form major viral proteins (S, M, N, E) [27]. Those viral proteins are processed through the endoplasmic-Golgi network to finally self-assemble to form a complete virion [28]. During infection, SARS-CoV-2 hijacks different host cell machinery, one such factor is the activation of Caspase 8, a key factor of extrinsic apoptosis that leads to host cell death [29].

2.4. Similarity with other viruses

Based on host type, coronaviruses can be classified into animal coronaviruses and human coronaviruses. The human coronaviruses can be divided into 4 different subfamilies: alpha, beta, gamma, and delta [3]. The subfamilies can further be classified into subgroups. Though the genome organization is similar but consecutive mutation at S protein and open reading frame (ORF) makes one coronavirus different from the other one [27]. Current SARS-CoV-2 has sequence similarity with SARS-CoV of 76% whereas BatCoV-RaTG13, the suspected precursor of SARS-CoV-2 has sequence closeness of around 97% [30].
The viral disease is highly contagious as transmitted by airborne, droplets, and touch-based [31]. Results of comparative analysis support SARS-CoV2 virus contains RNA as genetic material as other retroviruses. It is also found 91.2 % genetic similarity of nC0VID and Pan-SL-CoV which are found in pangolin of Guangdong province and 85.4 % with Pan-SL-CoV virus present in pangolins of Guangxi province [32]. The spike protein of the virus differs by only one amino acid of Spike protein [33]. Structural mutation at S protein led the SARS to detect host ACE2 receptor, while MERS recognize only DPP4 and SARS-CoV-2 detect ACE2 [28].

Moreover, regarding transmissibility, the Influenza virus shares a lot of similarities with SARS-CoV-2 [33]. Nevertheless, genomic analysis suggests MERS-CoV shares only 50 % of sequence similarity, the asymptomatic transmission of SARS-CoV-2 is very similar to MERS-CoV [32]. Structural mutation at S protein led the SARS to detect host ACE2 and CD147 receptor, while MERS recognize only DPP4 and SARS-CoV-2 detect ACE2 [28].

As shown in Fig. 3, the genome encoding for various nonstructural proteins, structural proteins (Spike, envelop, nucleocapsid and membrane) and accessories proteins are similar in the case of COVID-19, SARS, and MERS virus with minor differences.

2.5. COVID-19, a panic zone virus

SARS-CoV-2 symptoms bear a resemblance to the common cold and other viral infections and thusly, it masked itself and deceived medical professionals initially, maintaining a medium Reproduction rate of 2.25 and a medium Mortality rate of 5.7 % till June 7, 2020 [34]. Most importantly, the current novel coronavirus has never been seen before in humans which is why the human immune system was completely uneducated about it. It cannot draw public attention in the initial phase of a pandemic. In a comparison with Death Zone viruses like Ebola and smallpox, this virus has a very less mortality rate but a higher infection rate, and due to which it is spreading so quickly. In addition to that, asymptomatic transmission is another big issue, unwittingly patients are spreading the virus. Since it becomes a prime issue so quickly, generates a huge medical equipment supply shortage in the initial phase to fight against SARS-CoV 2. Until now, more than a hundred variants of SARS-CoV-2 have appeared to spread with high transmission efficiency [35]. Virologists announced recent coronavirus strain delta plus as a ‘variant of concern’ for its high transmission efficiency and great immune escape ability [35]. Thence, developing countries like India faced over 300,000 active cases in the month of May–June, affected by extreme oxygen shortage. Above that several life-threatening fungal variants are affecting coronavirus patients or patients who had it earlier. In between extreme challenges, now and then, it is very important to take COVID-19 protective measures very seriously.

3. Biomarker associated

The COVID-19 emergency makes the scientists to commit accelerating safe and productive vaccine development. Understanding the pathogenicity and particular biomarkers associated with SARS-CoV-2 benefitting as the knight in shining armor for vaccine development [142]. Moreover, reliable SARS-CoV-2 biomarker identification is required not only for covid-19 disease diagnosis but also to identify the pathogenicity level [142]. Biomarkers can also help doctors to classify patients in various risk groups based on pathogenicity of virus and disease progression to provide better treatment. Based on the pattern of Biomarker (Hematological, Biochemical, Inflammatory, and immune) identified in the patient, the severity of the disease can be also identified and monitored accordingly [25]. The biomarker found in the COVID-19 patient can be categorized as stated in Table 1. Biomarkers can be both quantitative or qualitative [36]. In the case of hematological biomarkers, the number of various blood cells is the indicator of the severity of disease but in the case of inflammatory markers (Interleukin, C reactive proteins) and biochemical biomarkers (Total bilirubin, Blood urea nitrogen, LDH, Creatinin, Cardiac troponin, Aminotransferase, Creatine kinase), both qualitative and quantitative parameters are important to understand the severity of the disease [25,35].

4. Type of diagnostic test for COVID-19

SARS-CoV-2 infected patients most commonly have a fever, dry cough, tiredness, dyspnea, etc.

In one study guan et al. found that 44 % of 1099 patients had a fever while 89 % developed fever before and after entering the hospital, cough (38 %), Sputum production in 34 %, and 19 % of patients with shortness of breath in China [20]. Because of, similar disease presentation many of the COVID-19 symptoms can be also be correlated with various other respiratory diseases such as influenza, SARS, and MERS [33]. Testing methods and tools used for COVID-19 diagnoses are also based on the type of samples isolated from the patient and the target molecule to be identified in samples (Fig. 5). If the sample taken is oropharyngeal and nasopharyngeal swab, in that case, the viral antigen can be detected by ELISA and LFA-based methods and the viral genome can be identified by PCR and sequencing method [36]. But if the sample is patient blood, then the diagnosis method will be restricted to blood cell counts, viral antigen, and antibody against the virus as shown in Fig. 4 [36]. Although, RT-PCR (Reverse transcription-polymerase chain reaction) exhibit a gold standard in COVID-19 diagnoses [38], other conventional COVID-19

| Biomarker's type | Up-regulated | Downregulated |
|------------------|--------------|---------------|
| Hematological    | WBC count    | T cell count  |
|                  | Neutrophil count | B cell count  |
|                  | Neutrophil - lymphocyte ratio | Platelet count |
|                  | Monocyte-lymphocyte ratio | Eosinophil count |
|                  | Erythrocyte sedimentation rate | Lymphocyte count |
|                  | Interleukin class (2, 6, 8, 10) | NK cell count |
| Inflammatory     | C reactive protein | Albumin |
|                  | Serum ferritin | Biochemical |
|                  | Total bilirubin | Blood urea nitrogen |
|                  | LDH | Creatinin |
|                  | Creatinin | Cardiac troponin |
|                  | Aminotransferase | Creatine kinase |
| Blood coagulators | D Dimer | Almandine |
|                  | Pro-thrombin time | Angiotensin II |
| Potential new    | Angiotensin II | Homocysteine |
|                  | Almandine | Angiotensin (1–9) |

Fig. 3. Representation of genome similarity from 5’UTR (Untranslated region) to 3’UTR (Untranslated region) of SARS-CoV-2 with SARS-CoV and MERS-CoV, where E, M, N represent envelop protein-coding gene, a membrane protein-coding gene, and nucleocapsid protein-coding gene respectively. The numbers like 3, 7, 8, 9, 10B, 13, 14 in COVID-19 genome, 3A, 3B, 6, 7, 7B, 8, 8A, 9B in SARS-CoV genome and 3, 4A, 4B, 5, 8 in MERS-CoV genome represent accessories protein coding region present in the genome. Reproduced from an open-access article [27] under the terms of Creative Commons CC BY.
diagnosis method includes immunological detection method like ELISA, examination of the pathophysiological condition of the affected organ by chest CT (Computed tomography) scan, NGS (Next-generation sequencing), etc [36].

4.1. CT scan

A CT or computed tomography scan permits doctors to examine the internal organs of the body. In, CT scan, a large number of chest x-ray are taken, and cross-sectional 3D images are developed, analyzed to detect abnormalities to verify the presence of COVID-19 infections [40]. Ground-glass opacity appears in the lower part of lobes and consolidation of the lungs [41]. In the early cases, small patchy pulmonary lesions appear. According to one report, a CT scan was found to be more sensitive than RT-PCR at a very early stage of infection [41]. It can be deduced from their findings that a CT scan conjointly with any other standard diagnosis technique can achieve excellent accuracy. CT scan not only helps to identify the infection due to COVID-19 (Fig. 6) but also helps to understand the condition of lungs and severity of disease, which guides the physicians to manage the patients related to infection [42].

4.1.1. Challenges with CT scan

CT has very low specificity (25 %) for COVID-19. The diagnostic imaging with CT overlaps with another disease like pneumonia [43]. Further, there is a considerable amount of uncertainty with CT in the case of asymptomatic patients [7]. In addition to that, repeated exposure to radiation can impose a biological impact in a long run [43]. CT system is a very expensive tool and requires an expert to operate and analyze the
data so, there is a need to develop other tools. The key challenges and advantages are mentioned in SWOT 1.

4.2. Nucleic acid-based test

4.2.1. Sequencing based

Sequence-specific detection of SARS-CoV-2 can be performed through nucleic acid sequencing, one of the highly specific molecular techniques to detect novel Coronavirus. This method is used to analyze the complete genetic material of the virus [1]. It is not only useful in diagnosis but also to evaluate the mutation pattern of the virus. Further, genomic analysis is very crucial in designing different molecular assays such as primer designing for PCR. Most of the current COVID-19 research uses Sanger sequencer and next-generation sequencing [44]. In the case of sanger sequencing, the patient sample is lysed for viral RNA isolation and some spike-in control RNA is mixed along with Viral RNA [45]. The spiked RNA is used for the normalization and quantification of the virus present in the sample [45]. A specific set of primers allowed the PCR amplification to facilitate the differentiation of SARS-CoV-2 viral RNA and spike in RNA upon electropherogram analysis [45]. On the other hand, most next-generation sequencing-based SARS-CoV-2 diagnoses mainly rely on Illumina sequencing and Oxford nanopore sequencing [44]. Unlike sanger sequencing, next-generation sequencing can target billions of target sequences simultaneously [44]. Genomic and sub genomic data analysis enables understanding the molecular features of pathogenesis and to develop a different diagnostic approach for accurate SARS-CoV-2 detection from a patient sample. It is reported that sanger sequencing can run 4000 tests/day with a single instrument which is 30 times faster than RT-PCR [46]. Understanding the different SARS-CoV-2 variants is very critical to follow the effectiveness of different vaccines. Moreover, it will be very tough to determine the viral mutation and infection progress without sequencing. The gold standard qRT-PCR can only be modified and adapted based upon sequencing. In addition to that sanger sequencing can deliver proper diagnosis at a 30 times faster rate than RT-PCR [44]. To test one sample with Sanger sequencing the running cost approximated to be less than 20$ [44]. The sequencing-related advantages and disadvantages are mentioned in SWOT 2.

4.2.2. PCR based detection

PCR is a molecular technology that amplifies the segment of the targeted genome present in the sample. RT-PCR continues to be a benchmark in COVID-19 diagnosis. As the global outbreak of COVID-19 is based on a newly discovered SARS virus, the handy diagnostic approach appears less. With the metagenomic RNA sequencing method, the proteogenomic composition of SARS-CoV-2 was found quickly on March 25, 2020. Then, the RNA Sequence (29870 bp) of COVID-19 was shared with the public and added to the GenBank sequence repository on Jan 10, 2020 [21]. After onwards, more than 1000 sequence was shared and added in the GISAID database from various scientific community all across the world [21]. Depending upon the complete genomic information, scientists developed an accurate diagnostic tool for targeting specific regions of the viral genome. SARS-CoV-2 is mainly a positive-sense single-stranded RNA virus, so the most appropriate method for diagnosis based on PCR is RT-PCR, where RNA is first converted to complementary DNA and then amplified for detection [21]. RT-PCR-based SARS-CoV-2 detection, mainly target, RNA dependent RNA polymerase gene (RdRp), ORF1ab, N, E, and S portion of genomic sequence [38]. Except for the RdRp, the sensitivity of other targeted genes is in a comparable range. While, for RdRp, slightly lower sensitivity is mostly caused by the reverse primer-template mismatches [38].

SARS-CoV-2 can be detected directly from collected biological fluid which has to be preprocessed for viral RNA isolation that needs to be converted to cDNA in presence of reverse transcriptase [47]. The converted cDNA of the virus is targeted now, and a segment of the viral genome is amplified with PCR [47]. Additional components (probe) are
also added to situate a foundation that hybridized with the complementary cDNA segment for amplification. The single-step Taqman probe allows real-time quantitative monitoring of the PCR cycle [43]. The result of the PCR amplification is analyzed based on control (positive control, negative control, and internal control) used in the reactions and the Ct value (Threshold value) obtained during amplification of the viral genome (Fig. 7). The COVID-19 positivity criteria depend on measuring the Ct value. The Ct value is the number of PCR cycles required to cross the threshold fluorescent signal value. In most cases of COVID-19 infection, the cut-off Ct value is between 35 and 40 [48]. A patient with a Ct value lower than 35 is considered as COVID-19 positive, whereas above 40 Ct value, clinically considered as negative [48]. A Higher Ct value represents a lower viral load in the sample [47]. Various FDA-approved RT-PCR kits are listed in Table 4 for SARS-CoV-2 detection.

4.2.2.1. Challenges with PCR. There are many possible causes of false-negative cases in PCR-based SARS-CoV-2 detection. Firstly, during the viral replication cycle sometimes RNA gets damaged due to lower stability, which leads to fragmentation of the viral genome in the bloodstream that imposes a major challenge for RT-PCR-based detection [47]. Secondly, the exact severity of lung infection and associated viral load cannot be measured by examining any type of body fluid [47]. Thirdly, PCR as a diagnostic tool requires benchtop size instruments, a large number of PCR reagents, long detection time, which limits its usage in under-equipped rural areas [2]. No test is 100% accurate. In one COVID-19 diagnosis-related study, the false-negative rate for qRT-PCR was found to be 9.3% while sensitivity was 90.7% [50]. Other than mentioned issues related to qRT-PCR, sample cross-talk, software problems may decrease the sensitivity. In another large-scale study in China, the false positive rate was found to be 30% of the total patients diagnosed [51]. The major advantages and drawbacks are presented in SWOT-3.

4.2.3. CRISPR based

Recent progress in CRISPR (Clustered regularly interspaced short palindromic repeats) research aside from revolutionizes genome engineering also opens up a new area for nucleic acid diagnostics-based pathogen detection. CRISPR is mainly a bacterial defense system against the infected virus. CRISPR RNA (crRNA) guides the CRISPR protein to destroy the target sequence that is attached with a complementary crRNA sequence [53]. Each CRISPR protein has its signature cleavage ability that makes one unique from the other. For instance, Cas9 binds with target dsDNA while Cas13a cleaves ssDNA. Cas14 can recognize ssDNA more efficiently than dsDNA [53]. The current, global pandemic escalates the interest to use CRISPR tool inaccurate point of care COVID-19 diagnosis.

Two highly renowned companies Mammoth Bioscience and Sherlock Bioscience are harnessing the power of CRISPR to develop a cost-effective, rapid, specific diagnostic tool for COVID-19 (Fig. 8). Mammoth bioscience is working on a CRISPR tool developed by Jennifer lab that uses the Cas12a enzyme for cleaving and degradation of ssRNA and fluorescent reporter activation [54]. Feng Zhang and his colleagues developed SHERLOCK system which uses Cas13 enzyme along with a programmed CRISPR system, which can target specific gene sequence for cleavage, degradation of target strand and activation of the fluorescent reporter [55]. In both systems when the guide RNA finds the specific sequence in the sample, it cleaves the identified sequence and gives fluorescence. These two CRISPR-based biosensors work is on the principle of optical biosensors, which will capture the fluorescence during COVID-19 detection [54,55]. Different CRISPR-based COVID-19 detection methods are summarized in Table 2. Further main merits and demerits of CRISPR related to SARS-CoV-2 detection are mentioned in SWOT-4.

4.3. Immunoassay

Apart from direct detection, SARS-CoV-2 can also be diagnosed indirectly considering the immune response of an infected individual. Nevertheless, recent research has shown serological detection of COVID-19 with different fluids including saliva samples [52]. Rather than nasopharyngeal swab, serological diagnosis mainly targets blood samples

Fig. 7. Reverse transcription quantitative polymerase chain reaction of nasopharyngeal swab is the conventional method of COVID-19 confirmation. (A) Typically, RNA is extracted from clinical sample to amplify with qRT-PCR. (B) Various ongoing research used qRT-PCR for clinical diagnosis without any pre sample treatment. The assay sensitivity is in comparable range in both the cases however average threshold cycle value (Ct) increases in direct testing [52]. Reproduced with the permission from the Elsevier.
It was found that blood enriched by the detectable concentration of antibodies within 7 days of infection in 50% of affected individuals and within 14 days for all individuals. This primary immune response detection is very important to inspect community transmission. Analysis of IgG and IgM are two main antibodies for detection. The bioanalytical method that relies on antigen-antibody-based interaction is termed immunoassay. Traditionally LFA and ELISA, are two main immunoassay-based serological diagnostic methods. Specific antigens or antibodies are targeted for COVID-19 diagnosis. The stage of infection is crucial in deciding the diagnostic approach for COVID-19 detection. According to various studies, a minimum of 5 days is required for sufficient viral load and a minimum of 7 days is required for antibody development in the system. However, antibody concentration may decline after 7 days of infection. Many immunoassay platforms have been developed so far for rapid COVID-19 detection.

4.3.1. Challenges with immunoassay

The viral load does not remain constant throughout the infection which hinders the sensitivity of the test [67]. Other coronaviridae family viruses share similarities with the COVID-19 virus that causes cross-reactivity during the test to give a false-positive result [67]. Even being a less time-consuming, cost-effective method, the immunoassay-based test is less sensitive than a molecular test like PCR [66]. SARS-CoV-2 detection-related advantages and disadvantages are listed in SWOT-5.

4.3.2. Lateral flow immunoassay

Lateral flow immunoassay (LFA) is a paper build qualitative analyte for virus detection [36]. It was found that blood enriched by the detectable concentration of antibodies within 7 days of infection in 50% of affected individuals and within 14 days for all individuals. This

![Fig. 8. (A) Schematic of SHERLOCK based COVID-19 detection. The point of care method follows three simple steps and can be finished within an hour. (i) An isothermal amplification of extracted RNA (ii) Detection with Cas13-crRNA complex (iii) Visual readout in a paper strip. (B) Schematic of DETECTR based SARS-CoV-2 detection. Extracted RNA from nasopharyngeal swab is amplified by RT-LAMP. Followed by Cas12 based cleavage of target sequence and visual detection on lateral flow reader. Reproduced from open-access article [65] under the terms of Creative Commons CC BY.](image-url)

| SN | Name     | Target Sequence                          | Cas  | LOD                  |
|----|----------|-----------------------------------------|------|----------------------|
| 1  | SHERLOCK | SARS-CoV-2 S & ORF1ab gene              | Cas13a | 10-100 copies/μl [56] |
| 2  | DETECTR  | SARS-CoV-2 N & E gene                   | Cas12a | 10 copies/μl [54]   |
| 3  | CREST    | SARS-CoV-2 N gene (N1,N2, N3 segment)   | Cas13a | 10 copies/μl [57]   |
| 4  | CASdetect| SARS-CoV-2 RNA dependent RNA polymerase | Cas12b | 1 x 104 copies/ml [58] |
| 5  | iSCAN    | SARS-CoV-2 N&E genes                    | Cas12a | 10 RNA copies/reaction [59] |
| 6  | ENHANCE  | SARS-CoV-2 N gene                       | Cas12a | 1.3 copies [60]     |
| 7  | AIOD-CRISPR | SARS-CoV-2 N gene                      | Cas12a | 1.3 copies [61]     |
| 8  | FELUDA   | SARS-CoV-2 N & NSP8                     | Cas9  | 10 copies/sample [62] |
| 9  | CRISPR-FDS | SARS-CoV-2 ORF1ab                      | Cas12a | 10^8 ampls/sample [63] |
| 10 | CONAN    | SARS-CoV-2 N gene                       | Cas3  | 10^8 copies/sample [64] |
detection technique that uses a coating of antigen or antibody on the surface of a strip which bound to either viral protein or antibody developed against the virus [68,69]. Some lateral flow immuno-assay also uses aptamers as bioreceptor. When the sample from the infected person having SARS-CoV-2 is placed on a strip coated with antibody, the antibody binds with COVID-19 protein and shows the color change on a paper strip [69]. Generally, LFA is a qualitative test. As indicated in Fig. 5, the sample is mounted at one end of the paper strip, preferentially on the sample pad. Due to capillary interaction, a sample containing target analyte migrates towards a conjugation pad, that is coated with the probe, specific against the target analyte of interest. Upon the formation of a conjugate complex, it migrates towards the detection pad. Some lines in the detection pad are there to monitor the antibody-antigen reactions and are termed as M line, where some act as a control for the test, and some for guidance (G line) as shown in Fig. 9 [68]. The particular analyte recognition on the detection pad produces color read-out to be detected with the naked eye. The efficiency of color read-out can be adjusted by using different nanoparticles like colloid gold modified probe on conjugation pad [70].

In comparison with the gold standard, PCR test, lateral flow immunoassay based serological detection require less complex technical setup, rapid detection, and cost-efficient [70]. While inefficacy of early infection detection limits the ‘immunity passport’ detection using LFA [13].

There are various Point of care based rapid diagnostic kits, based on LFA that have already been developed by several companies worldwide for COVID-19 detection as stated in Table 3 and 5.

4.3.3. ELISA immunoassay

ELISA-based IgM, IgG detection exhibited a very specific and sensitive detection of COVID-19 so far.

Enzyme-linked immunosorbent assay is commonly known as ELISA because in this process enzymes are conjugated with specific antibodies to detect various proteins and pathogens present in the system [72]. It can work either way whether to detect COVID-19 viral antigen or antibody developed against the COVID-19 inside the host body. This system uses microtiter plates containing 96 wells, where the antibody is coated or fixed at the surface and then the sample containing the specific analyte (virus, proteins, antigens) is added, the fixed antibody with the analyte conjugate is identified by enzyme tagged antibody in presence of specific substrate which produces color, fluorescence or luminescence that can be visually identified [72]. The same procedure is followed for antibody detection for COVID-19 [73,74]. Companies like BioRad and Euroimmun US Inc have already developed ELISA-based methods for IgG and IgM detection for COVID-19 infection diagnosis as stated in Tables 3 and 5.

4.3.3.1. Antigen based. The antigen-based detection technique is important for the diagnosis of COVID-19, as it gives the direct result of the presence of virus inside the system and also helps in early detection of
COVID-19 to prevent further infection [75]. There are multiple proteins like envelope, spike, nucleocapsid, matrix, and others associated with the COVID-19 virus, which can act as potential biomarkers for COVID-19 diagnoses [76]. The problem with antigen detection is, as the viral load reduces after the first week of infection, the amount of viral protein also reduces and becomes difficult to detect. C-reactive proteins and D-dimer along with other various biomarkers associated with COVID-19 used as promising antigens candidates for the diagnosis of this infection [37]. Cytokines which are also considered the inflammatory biomarker can also be also detected. Other than performing individual immunooassays with ELISA, the principle of ELISA can also be integrated with biosensors for rapid and sensitive diagnosis.

4.3.3.2. Antibody-based testing. Development of antibodies in the human host in case of COVID-19 infections has been shown after a week of the first symptom appear in the patient. A high quantity of IgA and IgM has been identified using recombinant viral protein within 3 days of first symptom development [77]. It is also found that the specificity for specific antibodies against COVID-19 in ELISA is greater than 80 % [78]. Spike protein 1 has shown promising results in binding COVID-19 antibodies [78]. Whereas the case of COVID-19 diagnoses with both ELISA and colloidal gold immunochromatographic kit shows 100 % sensitivity and similar specificity [77]. There are more than 100 kits available in the market for the detection of antibodies specific to COVID-19 in an emergency. This type of test is rapid and cost-effective in comparison to other molecular tools. The antibody-based test also helps in identifying the history of pre-exposure to the virus of the patient. These tests will be useful in the re-opening of social engagement where a strong immune response against the virus can be detected.

4.3.3.2.1. Challenge with an antibody test. Detection of antibodies will be only beneficial for those post-symptomatic patients who have developed antibodies against COVID-19, but it is not advised particularly when the patient is in critical condition. These tests showed lower sensitivity in comparison to the PCR test [36]. It has been also seen many false-negative cases in antibody testing against COVID-19, can be due to lower titer value of the antibody, homologous protein present in the sample, or low sensitivity of the instrument [77].

4.4. Point of care test

Combining the rapid detection of disease-causing agents, with a pandemic preparedness plan can strengthen the healthcare sector to check future pandemic outbreaks.

Table 3

A different source of Kit based on Immunoassay with their Target analyte and method of detection (EUA authorized FDA approved, European Union Conformity Marked) [79].

| Source | Principle | Target Antibody |
|--------|-----------|-----------------|
| Abbott diagnostic co Ltd | LFA | Immunoglobulin G, Immunoglobulin M |
| DiaSorin Inc | CIA | Immunoglobulin G |
| Eurosimman US Inc | ELISA | Immunoglobulin G, Immunoglobulin M |
| Healgen Scientific | LFA | Immunoglobulin G, Immunoglobulin M |
| Jiangsu Macro & Micro Test | Gold in form of colloid | Immunoglobulin G, Immunoglobulin M |
| Ortho Clinical | LFA | Immunoglobulin G, Immunoglobulin M |
| Quidel Corporation | LFA | Nucleopapsid protein |
| AutoBio Diagnostics | Immunology based | – |
| Siemens Healthcare Diagnostics Inc | Immunochemistry | Total Antibody |
| Vibrant America | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Clinical Laboratories | Chemiluminescence | Immunoglobulin G, Immunoglobulin M |
| Assure Tech | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Hangzhou Biotest Biotech Co., Ltd | LFA | Immunoglobulin G, Immunoglobulin M |
| Beijing Decombio Biotechnology | Immunology based | Immunoglobulin G, Immunoglobulin M |
| BIOSCIENCE | Immunology based | Immunoglobulin G, Immunoglobulin M |
| BNTX Inc | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Abbott Laboratories, Inc | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Cellex | Immunology based | Immunoglobulin G, Immunoglobulin M |
| ChemBio diagnostics | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Core Technology | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Roche Diagnostics | Immunology based | IL-6, Immunoglobulin G, Immunoglobulin M |
| PharmaTech | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Snibe Diagnostic | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Wadsworth Center | Microsphere based Immunology | Total Antibody |
| Nanoresearch | Immunology based | Immunoglobulin G, Immunoglobulin M |
| Mount Sinai Laboratory | Immunology based | Immunoglobulin G |

Table 4

List of nucleic acid-based diagnostic tools developed on the basis of emergency use authorized by FDA [79].

| Source | Mechanism of Detection | Site/antibody Target | Time limit | Detection limit | Point of care |
|--------|------------------------|----------------------|------------|----------------|--------------|
| Thermo Fisher | qReverse transcriptase -PCR | Open reading frame 1a, Nucleoprotein coding gene, Surface protein coding-gene | >2h | 10 copies/microliter | no |
| Roche | qReverse transcriptase -PCR | Open reading frame 1a | >2h | 9 copies/microliter | no |
| PerkinElmer | qReverse transcriptase -PCR | Open reading frame 1 ab, Nucleoprotein coding gene | >2h | 0.025 copies/ microliter | no |
| Sherlock Bioscience | CRISPR + LFA | Surface protein-coding gene, Open reading frame 1ab | 60 min | 10-100 copies/ microliter | yes |
| Cepheid | qReverse transcriptase -PCR | Nucleoprotein coding gene, Envelope protein coding-gene | 45 min | 0.25 copies/ microliter | yes |
| BioFire | qReverse transcriptase -PCR | Open reading frame 8, Open reading frame 1 ab | 50 min | 0.33 copies/ microliter | yes |
| Mesa Biotech | Reverse transcriptase -PCR + LFA | Nucleoprotein coding gene | 30 min | 200 copies/reaction | yes |
| Abbott | Isothermal nucletic acid amplification | RNA dependent RNA polymerase-gene | 5/13 min | 0.125 copies/ microliter | yes |
| CDC | qReverse transcriptase -PCR | Nucleoprotein coding gene | >2h | 3.2 copies/ microliter | no |
| Mammoth Bioscience | CRISPR + LFA | Envelope protein-coding gene, Nucleoprotein gene | 30 min | 70-300 copies/ microliter | yes |
In a situation like a pandemic, point of care testing (POC) is very crucial in terms of disease management [143]. The priority is to understand the pathogen before developing any suitable diagnostic tool. It allows rapid detection of analyte near to the patient that enables diagnosis at a very early stage and routine monitoring to improve treatment at the right time. Market analytics expect the POC diagnostic market to grow $81.37 billion by 2028 at a compound annual growth rate of 9.4% [80].

Analyzing the recent trend, we can definitely say that COVID-19 is the real game-changer in the POC diagnostic market. Since new technologies are coming very often in this COVID-19 situation, POC as a whole moves towards a crucial point in the diagnosis and healthcare sector. Tables 3 and 5 consist of such point-of-care-based tests, which are FDA approved for emergency use. These tools are either based on the detection of viral RNA or immunological components like antigen or antibodies.

5. Biosensors and their properties

Biosensors are analytical tools, a combination of organic (Biological), inorganic (transducer), and signal processing component, which can detect an analyte of interest and provide quantitative or qualitative information. It is a sensing tool that contains a biological component as the receptor and a physical element as the transducer, for the detection of a specific molecule [10]. It is generally composed of three important components which are Bioreceptors, electrochemical interference, and transducer [11]. Specific bioreceptor in a sensitive biosensor interacts with a particular analyte, while electrochemical interference provides a surface for reaction and helps in the generation of a biochemical signal into an electrical signal for effective detection [67]. An accurate, rapid biosensor must fulfill all WHO prescribed ASSURED criteria (Affordable, Sensitive, Specific, User-friendly, Rapid and Robust, Equipment-free, and Deliverable to end-users) to become a useful and unique tool [81].

5.1. Designing a biosensor

A marvelous advancement in the healthcare sector has been made right after the invention of the glucose biosensor [83]. Both in terms of innovation and application, biosensor has adapted many technical strategies to enlarge as micro size, cost-effective, sensitive, multifunctional, and powerful analytical device (Fig. 10) [87]. Current biosensor research very much emphasizes improving biosensor designing so to address the shortcomings [84]. Every biosensor has two parts as bio-elements and sensor elements. Bio-element acts as the receptor of Biosensors which determines the specificity of biosensors, whereas sensing elements are used in signal transduction in biosensors [85,86].

There are various methods proposed for combining bio-elements with sensor elements of biosensors, here only four methods of coupling are focused which are mostly used in the development of biosensors [86,87]. They can be as stated below.

a) Matrix entrapment
b) Membrane immobilization
c) covalent amalgamation
d) physical adsorption encapsulation

5.2. Classification of Biosensors

Biosensors can be divided into various classes based on multiple scales. Based on signals sensing method and transduction principles they can be classified as electrochemical, optical, radiation-sensitive, mass sensitive, and many more (Fig. 11) [88]. Based on the Biorecognition element prospective biosensors can be grouped as protein-based, ligand-based, enzyme-based, nucleic acid-based, saccharides-based, oligonucleotide-based, aptamers-based as shown in Fig. 7 [89]. The biosensor performance depends upon these immobilized components, to
5.2.1. Immunosensors

Immunosensors utilize the concept of biochemical immunoassay where mainly antibody-antigen interactions are identified on the surface of the transducer and an output signal is measured [1]. In this type of sensor, both monoclonal and polyclonal antibodies can be used which binds with the complementary antigen for detection of specific pathogens. Immunosensors are mainly classified into two broad categories: labeled and label-free immunosensor [70]. Various immunosensors have been developed by multiple companies on an immediate basis for COVID-19 diagnosis. Some are listed in Tables 3 and 5. An immunosensor measures the biochemical interaction between the target analyte and specific receptor through various detection methods like optical, electrochemical, fluorescence, or other detection methods [11]. Though label-based immunosensors can seriously improve the sensitivity and detection efficiency however wipe out of enzyme labels, purification, and washing steps increase the complexity and cost factor [91]. In addition to that nonspecific binding, stability of major concern [91]. Inadequacy to combine the sensing and detection area to a single platform is a serious problem in the commercialization of immunosensors.

5.2.2. Enzyme based biosensor

Enzymes are often used as biorecognition elements for biosensor detection. These immobilized enzymes are very specific to the analyte of interest and catalyze biochemical reactions for successful detection. The most efficacious commercially available biosensors are for evaluating blood glucose levels, developed by Guilbault [83]. Enzymatic biosensors are more stable, non-toxic, sensitive, and can easily be tagged with a radiolabeled and fluorescent molecule [92]. Different types of enzymes can be immobilized on the sensor surface. HRP (Horseradish Peroxide) is a commonly used enzyme in a sensor like this [92]. Various factors influence the performance of enzyme-based biosensors such as medium pH, temperature, cofactor, etc [93]. In affinity-based sensors, the utilization of enzymes is very common. ELISA-based kits developed by various companies mostly use enzymes as the sensing tool. Other than traditional ELISA, diagnosis of SARS-CoV-2 very much rely on enzyme-labeled antibodies that mediate signal generation and amplification for selective identification. Due to high catalytic efficiency and specificity towards analyte identification, enzyme-based biosensors conceived an extensive usage in biosensors [93]. The antibody-tagged specific enzyme has a set of substrates or particular substrates to catalyze and different detection methods can be engrossed for detection of the catalyzed product (Fig. 12). Such as in a recent publish electrochemical detection of SARS-CoV-2 spike (S) protein employed alkaline phosphatase enzyme tagged secondary antibody to catalyze 1 naphthyl phosphate substrate on the surface of a screen-printed carbon electrode (SPCE). [94]. However, due to the instability of enzymes depending on the environmental condition, nanotechnology stole the limelight in biosensor design [93,141]. Above all of that, the capacity to work with small sample volume, sensor robustness, and high substrate specificity will always employ enzymes as across-the-board sensing elements [93]. Various advantages and disadvantages of the enzyme-based biosensor are listed in SWOT 6 in consequence of COVID-19 diagnosis.

5.2.3. Optical biosensor

Due to simple, straightforward, cost-effective techniques, optical biosensors have the potential to become a major device in COVID-19 diagnosis [146].

These are the sensors in which labeled analyte excited with a laser and the signals are generated [96]. The first optical biosensor was based on fiber optics. The optical biosensor is divided into several types based on the light used and absorbent, UV–Visible sensor, surface plasmon resonance, FRET (Fluorescence Resonance Energy Transfer) sensors, and others are common optical sensors [11,96]. Changes in the intensity of light are detected by this type of sensor. LSPR (Localized surface plasmon resonance) based optical biosensors for SARS-CoV-2 detection have been developed where the viral genome is bound to gold nanoparticles and then illuminated with a light source as shown in Fig. 13 [96]. The heat generated is detected by LSPR detectors, the signals generated by the detector are used to analyze the presence of SARS-CoV-2 in the sample [97].

Melting temperature analysis is very crucial in qRT-PCR. The primer annealing temperature is usually set up slightly lower than melting temperature to stimulate successful primer binding. Qiu et al. followed the same idea to develop a plasmonic photothermal biosensor for Covid detection [98]. They thiol fabricated cDNA on the gold chip to immobilize the complementary SARS-CoV-2 RdRp sequence at a temperature rise of 41°C [98]. The photothermal energy generated on gold nanochip is further converted to plasmon energy and transduced for sensing. Based on the signal-to-target sample size relationship the LOD for this dual-functional biosensor was found to be 2.26 × 10⁴ copies [98]. An extensive study on sandwich assay-based optical detection has substantially developed the scientific research towards combating this deadly disease. Ramakrishna et al. have developed a fiber-optic biosensor for the detection of COVID-19. They immobilized the capture antibody on optical fiber and ultimately a sandwich was formed on the U bent fiber probe [99]. AuNPs were tagged as a plasmonic label for the evanescent wave absorbance sensing to diagnose the target analyte [99]. Positive and negative consequences related to SARS-CoV-2 detection are listed in SWOT-7.

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**Fig. 12.** Enzymatic biosensor for SARS-CoV detection. Enzymatic reaction upon aptamer-viral spike protein interaction useful for virus detection based on electrochemical measurement. Reproduced from open-access article [95] under the terms of Creative Commons CC BY.
5.2.4. FRET biosensor

Fluorescence resonance energy transfer (FRET) is one type of non-radiative energy transfer event between the donor and accepted fluorophore separated between a distance of 1–10 nm [100]. It is a very important analytical tool, particularly in the field of biological research to quantify in vivo protein to protein interaction [100]. High sensitivity towards the intermolecular separation between fluorophore pairs, makes the FRET biosensor a powerful analytical tool. In one experiment Weng et al. developed a sandwiched FRET biosensor for the detection of avian coronavirus. They used molybdenum disulfide (MoS2) fluorophore modified antibody to bind with target antigen while Alexa fluor 488 labeled antibodies to form a sandwich between MoS2-Ab and target analyte [101]. Based on intermolecular distance-dependent energy transfer between Alexa-fluor-488 and MoS2, they detected avian coronavirus in the linear range of $10^2$ to $10^6$ EID50 and limit of detection of $4.6 \times 10^2$ EID50 [101]. The development of a promising FRET design can certainly be a powerful device for SARS-CoV-2 detection. In another experiment, Brown et al. developed a FRET biosensor based on the proteolytic cleavage of two fluorogenic peptide molecules, fabricated on the sensor surface. Shown in Fig. 14 [102]. The peptide hydrolyzation leads to a notable increase in fluorescence signal. The increased signal is directly related to the concentration of SARS-CoV-2 extracellular protease. This extremely sensitive biosensor can achieve a LOD of 9.7 ± 3 pfu/mL [102]. The main advantages and drawbacks associated with the FRET biosensor are listed in SWOT-8.

5.2.5. Colorimetric biosensors

To address the COVID-19 problems, mass testing is very crucial. Considering that, the colorimetric test is very simple, and qualitative or quantitative results can be identified by naked eyes. Colorimetric biosensors are tools that work on the principle in which color change is detected associated with specific biochemical reactions.

Fig. 13. Localized surface plasmon resonance-based diagnosis of COVID-19 [97]. Reproduced with permission from Elsevier.

Fig. 14. Schematic of sensitive, fast FRET assay for SARS-CoV-2 detection. (A) Due to internal quenching of fluorogenic peptides screen high FRET signal. (B) Enzymatic hydrolysis of peptide bonds increases the intermolecular distances and eventually the FRET signal decreases. Reproduced from open-access article [102] under the terms of Creative Commons CC BY.
between analyte and sensing receptors on the biosensor surface [103]. Detection of color can be done easily without expensive tools [103]. Sensitivity can be also increased by using nanomaterials such as colloidal golds [104]. In the case of COVID-19 detection, mainly viral structural antigen is targeted with a specific antibody. When the reaction between the antigen (viral protein) and antibody takes place, the particular color on the strip of the biosensor is generated [104]. The color that appeared on the strip can be visualized by naked eyes or can be detected by detectors present in biosensors. Nanotechnology has marked tremendous advantages in colorimetric assays [141]. AuNP is very convenient because of its attractive optical properties, tunable surface plasmon, and exceptional photostability [105]. Recently, a colorimetric biosassay has been developed based on the detection of color change with the naked eye. The SARS-CoV-2 antisense N oligonucleotide tagged with AuNP that in presence of complementary interaction change its surface plasmon can be detected (Fig. 15). Further to visual detection, endonuclease enzyme RNase H was used to visually detect the precipitation of AuNP [106]. As given in Table 3, there are industries like Jiangsu Macro & Micro-test and others that have used colloidal gold to enhance the sensitivity of LFA-based colorimetric biosensors. Even having many advantages, colorimetric biosensors contain several disadvantages, listed in SWOT-9, as well.

5.2.6. Electrochemical biosensors

The idea of electrochemical biosensors mainly emerges from the classical discovery of glucose biosensors. An electrochemical advanced transduction system can detect electrochemical responses where the biochemical information is transduced to an electric signal [96]. High sensitivity, easy miniaturization, small volume requirement, low cost makes this an excellent choice for analyte detection [107]. Electrochemical sensors are such an analytical platform that can be used for both qualitative and quantitative analyte detection [107]. These sensors can further be classified into various groups based on the sensing method, amperometric, voltammetric, potentiometric [1]. Amperometric biosensors are commonly used sensing tools for virus and pathogen detection. It detects the change in current at constant potential during the electrochemical reaction [9]. This tool is highly sensitive, rapid-response time, easy to use, and cheaper than other tools. Potentiometric biosensors are a type of electrochemical sensor to detect potential differences, due to chemical reactions [9]. This type of sensor mostly uses enzymes as a biorecognition element that senses the hydrogen (Proton) ions that are released or absorbed during reaction causing the change in potential of the surrounding environment on sensors [107]. Whereas impedimetric electrochemical biosensors depend on the change in impedance on the sensor surface [9]. Mainly, recognition of target analyte by biorecognition element contributes to the change in impedance, which is detected by electroactive species [108].

Designing suitable electroactive material is very essential in the case of the electrochemical biosensor [145]. Electrically conductive solid materials are generally used such as gold, copper, carbon-based, nickel, etc. Further, advancement in nanotechnology manifested a huge significance in an electrochemical sensor. Nanoparticles modify the transducer part of a biosensor to make it more electrokinetically active [11]. As the knowledge of SARS-CoV-2 has increased, it opens several new ways of accurate detection. A hypothetical schematic of an electrochemical biosensor is exhibited in Fig. 16 for the successful detection of SARS-CoV-2 on the graphene surface. A novel sensor based on the fabrication of AuNP on the surface of fluorine-loaded tin oxide doped electrodes has been successfully developed for SARS-CoV-2 detection [109]. They mainly immobilized antibodies against SARS-CoV-2 spike protein. In the quest for rapid diagnosis, this biosensor can accurately detect SARS-CoV-2 S protein within 10–30 s [109]. Though antibodies are very specific towards target analyte, in terms of stability, immobilization efficiency, aptamer shows superiority over antibodies. In one experiment, Tian et al. developed a dual aptamer-based detection of SARS-CoV-2 N protein on the gold electrode surface. They used Au@Pt/MIL-53 as multifunctional catalysts for the generation of the electrochemical signal [110]. This electrochemical aptasensor can detect 8.33 pg/ml of viral particles in real-time [110]. Here, we have listed some of the electrochemical attempts to detect SARS-CoV-2 in Table 6. The advantages and limitations of SARS-COV-2 specific electrochemical biosensors are listed in SWOT-10.

5.2.7. Piezoelectric biosensors

A piezoelectric biosensor is a type of label-free sensor that works on change in oscillation frequency upon biorecognition event [118]. These types of biosensors are based on piezoelectric materials, that can generate voltage when mechanically stressed. Such a sensitive sensor can even sense a very low femtogram of mass [118]. The surface of these piezoelectric crystals coated with receptors which on binding with the analyte, reduces the oscillation frequency of quartz-crystal material (Fig. 17) [117]. It is also used in the diagnosis of pathogenic microorganisms from various clinical samples [117]. In this current global pandemic situation, SARS-CoV-2 has spread almost everywhere. In such a scenario, the piezoelectric biosensor is an alluring device as it doesn’t need any power source.

5.2.8. Magnetoelastic biosensor

Magnetic biosensors start to get some particular attention in the last few years.

These sensors are made from a ferromagnetic amorphous alloy. Wireless Sensors use a coil system where the signal is generated away from the coil and sensing is done remotely when the sample is subjected to the sensor [117]. The change in magnetic field causes magnetoelastic resonance, which is detected by a non-contact signal collector coil, [21]. Pietschmann et al. have developed a magnetic SARS-CoV-2 antibody detection set up based on immobilization of S protein on the sensor surface. They found the limit of detection of 3.36 ng/ml in the serum sample while 2.96 ng/ml in PBS [119]. Even so, having several advantages, magnetic immunosensors are usually ignored when it comes to POC diagnostics. One main disadvantage, associated with the magnetoelastic sensor is the dependence of magnetic labels for detection [117]. Nevertheless, scientists are trying to optimize that part so to make a quality POC market for magnetoelastic biosensors.

5.2.9. FET biosensor

The enzyme-electrode biochemical-based biosensor was first developed by Clark and Lyons in the year 1962 [147]. After that, several types
was developed by Bergveld approximately third gate electrode through an insulator is called a FET semiconductor between the source and drain terminals is regulated by a permission from the Elsevier. (For interpretation of the references to color in this attachment of SARS-CoV-2 (f) on incubation of clinical sample (e) electrochemical set up and signal processing unit for detection (g) [116]. Reproduced with the execution of the references to color in this figure legend, the reader is referred to the Web version of this article.)

Fig. 16. Schematic illustration of designing of electrochemical immunosensor for SARS-CoV-2 detection. Deposition of graphene (a) and gold nanoparticle (b) on electrode surface increases overall surface area and conductivity of sensor surface (c). Immobilization of anti-SARS-CoV-2 spike antibody (d) on sensor surface for the target is negatively charged. The reverse will be applicable for a p-type n-type and p-type, where electrons and holes are the charge carriers, respectively. In the case of a positively charged target molecule, there will be a drastic increase in conductance in the n-type sensor due to the structure [152].

Table 6
Various electrochemical biosensors for the detection of SARS-CoV-2.

| SN | Target | Recognition element | Detection method | LOD |
|----|--------|---------------------|------------------|-----|
| 1  | S Protein | SpAb-AuNPs-FDTO | DPV | 90 fM [112] |
| 2  | S Protein | G-PBASE-anti-S Ab | FET based detection DPV | 242 copies/ml [113] |
| 3  | N protein | PdPd-Au printed board | DPV | 27 fM [114] |
| 4  | S & N Protein | Sandwich assay on SPE | DPV | 8 fM(N protein) [94] |
| 5  | N protein | Anti N Ab on SPE | SWV | 0.8 pg/ml [115] |
| 6  | S Protein | SpAb-PBASE-Graphene | FET | 16 pfu/ml [112] |
| 7  | N gene | PCR | DPV | 10 pg/ml [111] |

of sensing mechanisms have been developed. Among these, a unique kind of sensing approach was discovered, i.e. FET-based biosensor because of rapid growth in solid-state technologies [148]. Field-effect transistor (FET) biosensors grew to be one of the most promising alternatives due to their advantages like being fast, low-cost, and simple. At the same time, the ability to make highly sensitive and instantaneous measurements using small amounts of analytes makes them a suitable candidate for use in point-of-care testing and diagnosis [149]. The very first FET biosensor was developed by Bergveld approximately fifty years ago and evolving since 1970 in different forms and is an ideal approach for swift and precise detection of various analytes [150].

To define, “a solid-state device in which the electroconductivity of the semiconductor between the source and drain terminals is regulated by a third gate electrode through an insulator is called a FET” [151]. Generally, there are two kinds of FETs based on charge carriers namely n-type and p-type, where electrons and holes are the charge carriers, respectively. In the case of a positively charged target molecule, there will be a drastic increase in conductance in the n-type sensor due to electron aggregation. In contrast, the conductance will decrease if the target is negatively charged. The reverse will be applicable for a p-type biosensor. To enhance the sensitivity, nano materials are deposited on the ability to make highly sensitive and instantaneous measurements using small amounts of analytes makes them a suitable candidate for use in point-of-care testing and diagnosis [150].

5.2.10. Microfluidic biosensor
Microfluidics refers to the technology for the fabrication of micro-miniaturized devices having micrometer-sized channels and chambers, in order to allow small volumes of fluids to flow [155,156]. In recent years, there has been a tremendous increase in the integration of biosensing with microfluidics. Advantages like small sample volumes, rapid turnaround times, ease of multiplexing, and high portability offered by microfluidics, which make it suitable for biosensing applications. Precise control of droplets gives rise to miniaturization and rapid processing of samples for the application of lab-on-a-chip diagnostics [157]. These microfluidic-based point-of-care (PoC) devices can detect analytes and may provide rapid diagnostics even in remote regions and near a patient with limited-resource or non-existing healthcare settings [158]. This happened because of the integration of POC devices with smartphones and wireless communications. The selection of materials for fabrication should be such that, it should be optically transparent, easily modifiable and mechanically strong, and thermally stable. A single material cannot fulfill all these requirements.

A unique sensing scheme was developed by Liljehoj and group [159] which utilizes dually-labeled magnetic nanobeads for immunomagnetic enrichment and signal amplification on a microfluidic chip to detect SARS-CoV-2 nucleocapsid protein (Fig. 18). It provides advantages of enhanced detection sensitivity, minimal sample, reagent consumption, and simplified sample handling. It could detect concentrations as low as 50 pg/mL in whole serum within 1 h and 10 pg/mL in 5 × diluted serum.
Fig. 17. An overview on main approach for SARS-CoV-2 detection. (a) Cantilever as resonant sensor (piezoelectric material) detect the target viral antigen. Due to adsorption of viral antigen on fabricated antibody the mass changes lead to cantilever bending. That changes the signal. (b) voltage against time (c) amplitude against frequency. Reproduced from open-access article [117] under the terms of Creative Commons CC BY.

Fig. 18. Schematic illustrations of (A) microfluidic immunosensor chip highlighting the magnetic concentration of DMBs to the sensor surface, (B) microfluidic immunosensor chip for the smartphone-based diagnostic device, and (C) experimental setup and electrochemical sensing scheme using the PalmSens4-based sensing platform. Reprinted with permission from (Li, J.; Lillehoj, P. B. Microfluidic Magneto Immunosensor for Rapid, High Sensitivity Measurements of SARS-CoV-2 Nucleocapsid Protein in Serum. ACS Sensors 2021, 6 (3), 1270–1278). Copyright (2021), American Chemical Society.
within half an hour. To add to its portability, the chip was connected to a smartphone using a smart potentiotstat, which does not require external power sources.

Hwang-soo et al. [160] proposed a different procedure to detect SARS-CoV-2 using isothermal amplification of nucleic acids by utilizing a mesh having multiple pores. Blockage of pore takes place by the formation of DNA hydrogel when hybridization of pathogen and probe DNA occurs. It has a limit of detection (LOD) of 0.7 aM at 15-min incubation for SARS-CoV-2. But, it is necessary to optimize the mesh material, pore size, and precision microfluidics for achieving such value.

6. Overview of biosensing platforms for SARS-CoV-2 detection

In view of supply chain issues related to testing reagents and sample transportation, the current era imposes a serious emergency condition for global supplies. The US Centers for Disease Control and Prevention (CDC) has updated the guidelines for clinical sample collection and transportation. CDC recommends the clinical sample to store at 2–8 °C for 72 h and afterward sample storage in particular viral transportation media at – 70 °C [121]. These guidelines impose a huge difficulty to reach mass testing. Portable, automatic point of care devices eliminate the requirement for sample storage and transportation. Various smart POCT devices not only allow fast data acquisition but facilitate at-home diagnosis [144]. The gold standard qRT-PCR test prices Rs 2500–3200 ($42.96) depending upon various states of India while the accurate paper-based ‘FELUDA’ costs only Rs 500 ($6.80) [120]. Further scientists from IIT-Bombay have developed a printed circuit board (PCB) electrode-based electrochemical detection of SARS-CoV-2 with RS 40 ($0.05) [111]. Considering the current scenario, ICMR has approved a rapid antigen test which costs Rs 250 ($3.32) and provides a result within 15 min [122]. In comparison with RT-PCR that even takes 3–4 days to get the result, the POCT platform is very fast and has opened the opportunity for quick mass diagnosis [143].

Further advancement of nano biosensors can exponentiate remaining detection-related concerns. Other than the decoration of sensor surface with nanoparticles, discussed above, polymer-coated biosensors were found to be very convenient for fast, accurate detection [11]. Polymers containing acrylic groups are mainly favored for biosensor fabrication.

Above all mentioned methods of SARS-CoV-2 LFA biosensors is the key player with a view to the commercial market. After the integration of powerful CRISPR-Cas technique with LFA has been in the focus of public attention. Such biosensor not only reveals high specificity and sensitivity but being a low cost and on-site diagnostic tool, a non-specialist person can perform the test in real-time. As a point of care, diagnostics can’t be standalone with 100% accuracy, it is better to run such a rapid antigen test, combined with gold-standard RT-PCR. To combat concerned issues, researchers are going to develop a ‘sample in-result out’ POC nucleic acid detection system. Such as IIT- Kharagpur scientists develop an isothermal nucleic acid testing-based SARS-CoV-2 detection platform, called COV-IRAP, where the result can be interpreted with a free smartphone app [123]. To accelerate the detection and to come down with a more economical platform, microfluidics also joins the race. Researchers are trying to reduce the intricacy of PCR temperature control systems, with the help of a microfluidics system [124]. Here we have discussed some recent biosensors important for SARS-CoV-2 detection in Table 7.

7. Futuristic strategy for COVID-19 detection

There is a large number of kits and tools already present in the market but due to the lack of suitable properties as run time, cost, and complexity of technique it is important to develop more advanced POC tools. Among newly diagnostic tools developed for nucleic acid identification of COVID-19, LAMP (Loop mediated isothermal amplification) shows promising results compared to RT-PCR. Where PCR requires a sophisticated tool and facilities, this can be done at a constant temperature at 40-65-degree Celsius in the water bath [133].

RT-LAMP is based on the same principle where RNA of COVID-19 viruses is converted in cDNA and then amplified using 4 to 6 primers. LAMP is also integrated by Zhang's group [134] and Chiu's group [135] with CRISPR technology for the detection of COVID-19 viruses with the sensitivity of 10 copies/mL in POC format.

Another approach to develop a diagnostic kit can be based on Rolling Circle Amplification (RCA) [78]. This type of genome amplification method is generally found in viruses, which is also an isothermal method for nucleic acid identification in the sample. In this process, DNA is first circularized and then amplified with the high processive polymerase to achieve higher sensitivity. This method is used to develop a kit for COVID-19 diagnoses with higher sensitivity [21].

Lateral Flow-based assay for nucleic acid identification based on isothermal amplification can be an innovative option in developing diagnosis tools. There is a paper-based lateral flow assay (LFA) which have gain people's trust and popularity in the field of COVID-19 diagnostics [136]. Paper plastic-based LFA has been developed for malaria which uses folded paper for genome isolation and LAMP for genome amplification. 2D paper network-based LFAδs have been also developed for COVID-19 detections.

The major problem associated with the immune-associated LFA is having less sensitivity [136]. It can be solved by various approaches some

| S.N | Detection method | Material | Analyte | Detection time | LOD | Detection range |
|-----|------------------|----------|---------|----------------|-----|----------------|
| 1   | Magnetosensor    | Glossy carbon electrodes and magnetic beads | Viral S/N protein | 30 min | 8 ng/ml for S protein | [94] |
| 2   | Electrochemical sensor | Gold electrode modified with Fe3O4 | RNA genome | Real time | 200 copies/ml | [125] |
| 4   | Optical sensor | Optical fiber and gold nanoparticle modified protein | N protein | – | 1 pg/ml | [126] |
| 5   | Field effect transistor | Graphene layer | S proteins | Real time | 1 fg/ml | [113] |
| 6   | Breath sensor | Anl prism 700 instrument | N protein | 10 s | – | [127] |
| 7   | Piezoelectric immunosensor | Quartz crystal and magnetic particle | Helicase protein | 1 min | 3.5 ng/ml | [128] |
| 8   | Laser scanning microscopy (Optical sensor) | Quantum dot fabricated aptamer | N protein | – | 0.1 pg/ml | [129] |
| 9   | SPR biosensor | Gold chip/aptamer | RNA genome | Real time | 2 nM | 1nM/1 pM | [130] |
| 10  | Paper based sensor | COVID-19 ePad | Antibody | 45 min | 1 pg/ml | [131] |
| 11  | SPR based colorimetric sensor | Oligonucleotide tagged gold nanoparticle | RNA genome | 10 min | 3.6–3.9 copies/ml | [105] |
| 12  | Electrochemical biosensor | Reduced graphene oxide nanoflakes | Antibody | Within minutes | 2.8 × 10^15 M (Spine protein) 16.9 × 10^15 M (Receptor binding protein antibodies) | [132] |
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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