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Strategies and perspectives to develop SARS-CoV-2 detection methods and diagnostics

Rekha Jalandra\textsuperscript{a,d,1}, Amit K. Yadav\textsuperscript{b,1}, Damini Verma\textsuperscript{c}, Nishu Dalal\textsuperscript{a,e}, Minakshi Sharma\textsuperscript{d}, Rajeev Singh\textsuperscript{e}, Anil Kumar\textsuperscript{a,b,⁎}, Pratima R. Solanki\textsuperscript{b,⁎}

\textsuperscript{a} Gene Regulation Laboratory, National Institute of Immunology, New Delhi, 110067, India
\textsuperscript{b} Special Centre for Nanoscience, Jawaharlal Nehru University, New Delhi, 110067, India
\textsuperscript{c} Amity Institute of Applied Sciences, Amity University, Uttar Pradesh, 201313, India
\textsuperscript{d} Department of Zoology, Maharshi Dayanand University, Rohtak, 124001, India
\textsuperscript{e} Department of Environmental Science, Satyawati College, Delhi University, New Delhi, 110052, India

\begin{abstract}
To develop diagnostics and detection methods, current research is focussed on targeting the detection of coronavirus based on its RNA. Besides the RNA target, research reports are coming to develop diagnostics by targeting structure and other parts of coronavirus. PCR based detection system is widely used and various improvements in the PCR based detection system can be seen in the recent research reports. This review will discuss multiple detection methods for coronavirus for developing appropriate, reliable, and fast alternative techniques. Considering the current scenario of COVID-19 diagnostics around the world and an urgent need for the development of reliable and cheap diagnostic, various techniques based on CRISPR technology, antibody, MIP, LAMP, microarray, etc. should be discussed and tried.
\end{abstract}

1. Introduction

The COVID-19 pandemic caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has affected almost every country with 7,410,510 total cases and 418,294 deaths globally as of 12 June 2020 [1]. Since this virus is new and due to lack of any approved drug or vaccine, there is urgent need for a highly specific and sensitive diagnostic measures to identify infected people and isolate them to avoid the further spread of the virus [2,3]. Testing is paramount in the fight to slow down and minimize the spread of the virus. Effective diagnostic approaches need to be discussed and tried to detect the virus itself or the response of the host body to the virus. Currently, numerous diagnostic technologies are available for SARS-CoV-2 detection, and many are under development [4]. Recently, the World Health Organization (WHO) has enumerated the first two diagnostic tests for emergency use to detect COVID-19 to increase the accurate assessment of disease. The tests are based on the gene regulation polymerase chain reaction (RT-PCR) Corona virus and Cobas SARS-CoV-2, a qualitative assay for use on the Cobas® 6800/8800 Systems, and are for in-vitro diagnosis of COVID-19 [5]. Currently, the most popular diagnostic tool that identifies the viral RNA through amplification is RT-PCR. It is a sensitive technique that requires a small amount of input RNA but takes hours to get results [6]. Other tools that target virus includes microarray relying on the binding of the viral genome-specific probe and CRISPR technology that recruits Cas12/13 enzyme specific for viral genes for detection of SARS-CoV-2 [7,8]. As SARS – COV-2 infected individuals produce antibodies against the virus, these antibodies can be used for the identification of infected individuals. Antibody-based diagnostics give a positive result if they find binding antibodies that might have generated earlier during the infection and would serve to provide protection against reinfection thereby, limits their use in the detection of active diseases [9]. The sensors are another approach that presents low-cost promising diagnostic tools with high sensitivity to detect COVID-19 [10]. Most of the diagnostic techniques used now-a-days are time-consuming and require skilled expertise, but for large-scale screening, we need to develop more efficient methods that can be used for point-of-care (POC) detection to identify COVID-19 positive patients. Besides several diagnostic kits, US Food, and drug administration (USFDA) approved the first at home COVID-19 test kit with a home collection option to increase COVID-19 testing capacity [11]. In this crisis, where a collective pool of knowledge is a prerequisite, here, we recapitulate available updates on diagnostic methods such as PCR, microarray, molecularly imprinted...
polymer (MIP)-based sensor, CRISPR, etc. for COVID-19 detection.

2. Structure and genomics of SARS-CoV-2 are a vital premise for diagnostic development

As current and future COVID-19 detection methods are based on genomics and structure of SARS-CoV-2, it is pertinent to review the recent progress on these aspects. SARS-CoV-2, a novel coronavirus species, placed under the betacoronavirus genus based on genomic similarity and phylogenetic relationship with SARS-CoV (Fig. 1). Genome sequence of SARS-CoV-2 has 88% similarity with SARS-like bat derived coronaviruses, SL-CoVZC45, and SL-CoVZXC21. Among different coronaviruses, RNA-dependent RNA polymerase (RdRp) gene sequence is an extremely conserved sequence. According to the International Committee on Taxonomy of Viruses criteria, if a species shows less than 99.9% similarity for conserved RdRp sequence, it would be considered as novel species. RdRp sequence of isolated strain in Wuhan, China exhibits 86% similarity with existing SL-CoVZC45 coronavirus; therefore, CoVs were declared as a new species (SARS-CoV-2). SARS-CoV-2 has a single-stranded positive helix RNA genome of 30 kb with a GC content of 38% [12]. Whole-genome sequencing showed that the virus genome from different parts of the world exhibited sequence homology of more than 99.9% with SARS-CoV-2 isolated from Wuhan, China [13]. Homology modeling showed that the receptor-binding domain of SARS-CoV-2 and SARS-CoV differs only in a few amino acid residues [14]. The genome of SARS-CoV-2 consists of an open reading frame (ORF) 1a/b-coding region and four protein-coding regions flanking with the non-coding region on both sides. Starting from 5′ end protein-coding region, an s-region coding for spike protein, e-region coding for envelope protein, m-region coding for a membrane protein, and n-region coding for nucleocapsid protein are present [15]. Structural and accessory proteins (S, M, E, N-proteins) are translated from sgRNAs (single guide RNAs). The most abundant structural protein in coronavirus is membrane glycoprotein (~25–30 kDa), spans the lipid membrane thriced with the N-terminal domain on the outside and C-terminal domain inside the virion. S-protein (~150 kDa) recognizes and binds to the receptor present on the host cell, thereby responsible for viral infectivity. Scanning electron micrograph of the virus revealed that it is oval or spherical with stalk-like projections ending in round structure (spike) like other viruses of coronaviridae family. Spikes are essential for viral infectivity and host specificity. While invading host cell, furin-like proteases cleave S-protein into two parts: a receptor binding unit (S1) and a membrane-anchored fusion unit (S2). Envelope protein (8–12 kDa) determines the formation and composition of the viral membrane. Nucleocapsid protein protects and enfolds the viral RNA [16] (Fig. 2). SARS-CoV-2 binds to receptors on the cell surface via receptor-binding domain (RBD) present in their S1 subunit. RBD of SARS-CoV-2 is an almost identical 3-D structure with that of SARS-CoV and 76.47% amino acid sequence similarity, which uses spike proteins to bind with Angiotensin-Converting Enzyme 2 (ACE2) on host cell [17]. Thereby, it is believed that SARS-CoV-2 also enters cells by binding spike proteins to ACE2. SARS-CoV-2 contains ORF3 and whole ORF8 gene regions, which are characteristic features of bat-origin coronaviruses [12]. Scanning electron micrograph revealed that virus particle size ranges from 70–90 nm and invades various intracellular organelles, especially vesicles [13]. Immunofluorescent assays of the culture of Vero cells showing cytopathic effect with the convalescent serum from patients showed green signals in the cytoplasm; in contrast, no signal was detected in control serum. Though viruses recruit error-prone RNA polymerase for replication, so it is more likely to get mutated [18]. The emergence of new strains is a consequence of mutation. Mutation in SARS-CoV-2 enabled it to spread more efficiently from animal to humans and then from human to human. Mutation in ORF8 region at 28,144 position is stable and increases population frequency from 0 to 29% as the epidemic progressed (Fig. 3). These mutations became fixed during the outbreak and increase population frequency gradually. The phylogenetic analysis shows SARS-CoV-2 is the mutant product of SARS-CoV. The phylogeny indicates that the initial emergence of novel strain in Wuhan, China gained transmission between humans and was followed by worldwide outbreak transmission (Fig. 4). All the research reports on genomics and structure of SARS-CoV-2 are essential, which can become the premise of the development of more accurate, sensitive, and effective detection methods.

3. PCR-based detection

Due to the increasing incidence of SARS-CoV-2, PCR based detection is routinely used for the diagnosis of COVID-19. RT-PCR or qPCR is a molecular biology technique used to study gene expression at the
transcript level. It involves the following steps: RNA isolation from samples and cDNA synthesis using reverse transcription kit. PCR was set by making a working mix of the buffer, dNTPs, primers of the target gene, Taq polymerase, cDNA template and, SYBR green dye. PCR mix incubated in a PCR machine and fluorescence generated by PCR amplification is measured by machine to give Cycle threshold (Ct) values. Ct values of control and experimental samples are compared, and relative expression is estimated (Fig. 5). On 14 January 2020, the protocol of RT-PCR for detection of 2019-nCoV was published on the WHO website for its use of COVID-19 detection [19]. Other than regularly used RT-PCR for diagnosis of COVID-19, some other studies used other diagnostic techniques like RT-insulated isothermal PCR (RT-iiPCR) [20], reverse transcription loop-mediated isothermal amplification (RT-LAMP) [21] and a one-step rRT-PCR assay [22] in which unique TaqMan probes is used. RT-LAMP has a high specificity for detection of MERS-CoV, and its sensitivity is comparable to RT-PCR. RT-iiPCR and a one-step rRT-PCR assay also show similar sensitivity and specificity, but RT-PCR is used mainly for the diagnosis of MERS-CoV. The selection of specimen type and time of taking that sample for RT-PCR is also a crucial point for the diagnosis of SARS-CoV-2.

The PCR-based first test for diagnosis of SARS-CoV-2 was constructed in Germany by Corman et al., at first, before the release of sequence, RT-PCR assay was designed following SARS or SARS-like coronavirus because it was supposed that 2019-nCoV is SARS-related.

Fig. 2. Schematic of structure of SARS-CoV-2.
Diagram showing the single stranded RNA genome and proteins present in coronavirus.

Fig. 3. Heatmap showing the mutation in ORF 8.
The trend of the mutation sites over time is displayed in the form of heatmap. Population frequency of site 28,144 located in gene ORF8 slowly raised from 0 to 35.98 %, and then dropped to 16.96 % as the epidemic progressed (see the area that gradually changes from blue to red). Image has been taken from http://bigd.big.ac.cn/ncov/variation/heatmap.
After getting the sequence information, the RT-PCR assay was designed according to sequence similarity with SARS-CoV-2 after sequence alignment. Corman et al., developed a detection method that can differentiate between SARS-CoV and SARS-CoV-2. At first, three assays were done: E gene assay, N gene assay, and RdRp gene assay. But due to less sensitivity of N gene assay, only two assays were selected where E gene assay works as a first-line screening tool and RdRp gene assay was done for confirmation of this testing. They used two different probes: one binds with 2019-nCoV and SARS-CoV and 2nd one bind only with 2019-nCoV. Results show that 2019-nCoV related probe only binds with 2019-nCoV but not with SARS-CoV, so these assays are highly sensitive and specific [23].

Chu et al., designed two 1-step quantitative real-time reverse transcriptase PCR assays which detect two different regions: ORF1b and N as these are the highly conserved sequence in sarbecoviruses. Throat swab and sputum samples were tested from two patients along with positive and negative controls, and results are highly specific.

Fig. 4. Tree showing genomes of SARS–CoV-2 worldwide. Phylogenetic tree shows the 4645 genomes sampled from December 2019 to April 2020 in different geographical regions. Each color represent specific country where genome has been sequenced. Image has been taken from GISAID database (https://www.gisaid.org/epiflu-applications/next-hcov-19-app/).

Fig. 5. Diagnosis of COVID-19 using PCR.
a) Respiratory sample collection from patient b) Extraction of RNA from sample c) Reverse transcription for cDNA synthesis d) DNA amplification using RT-PCR e) COVID-19 positive DNA sample shows amplification, if sample is negative for COVID-19 then it doesn't show any amplification.
and sensitive. N gene RT-PCR works as a screening tool, and ORF1b
used as confirmatory testing [24]. Konrad et al., done laboratory testing
for comparing RT-PCR assays: QuantiTect Virus + Rox Vial kit
(QIAGEN), SuperScript III One-Step RT-PCR Platinum TaqDNA Poly-
merase (Invitrogen, Darmstadt, Germany) and a commercial kit: Real-
Star SARS-CoV-2 RT-PCR kit 1.0 (Altona).

The protocol and thermal conditions of each assay were adjusted
according to the manufacturer, but the concentration of probes was
taken the same as published before by Corman et al., in 2020. These
assays show unspecific E gene signals but using SuperScript III these
unspecific signals reduce to 5%, so SuperScript III RT-PCR is better as
given by Corman et al. This research also recommended RealStar SARS-
CoV-2 RT-PCR kit 1.0 (Altona) as it does not give any unspecific E gene
signals [25]. In a study, a combined approach was developed by using
RT-PCR, CRISPER- based assay, and metagenomic next-generation se-
quencing (mNGS) for diagnosis of pneumonia patient, which later finds
positive according to RT-PCR and CRISPER-based assay [26]. So, by
combining these approaches, equitable diagnostic results can be found.
An article published in a magazine named Semiconductor Digest stated
that Cepheid, a California company, made a new point of care detection
test: Xpert® XpressSARS-CoV-2. This test can give results only in
45 min, and it requires less than one min for sample preparation, so
even healthcare workers can use it easily [27], POC devices are easy to
use and less time taking so these can be used primarily for diagnostic
purposes in the next few months.

4. Antibody-based detection

Antibodies (Abs), also called immunoglobulins, are Y-shaped pro-
teins produced by the immune system when a foreign substance (like a
virus) enters the system, and then these molecules bind to the foreign
substance and neutralize it. Simply put, they are the weapon used by
the immune system to fight new infections [28]. The SARS-CoV-2 is
being tested for an approach to treating new diseases that worked for
the Ebola and Zika viruses, which involves using antibodies to the
disease as a drug. Because of exquisitely affinity, target specificity,
designed and engineered nature, they have played a significant role in
the number of diagnostic approaches [29]. Currently, a variety of
COVID-19 tests are already available that determine the viral genome
from mucosal membrane swabs, till now, eleven POC diagnostics for
COVID-19 are reported, which is mainly based on molecular and anti-
body-based tests. Various research groups and companies are trying
hard to develop antibody tests which include lateral flow immu-
noassays (BioMedics rapid test and Surescreen rapid test case-
tte), time-resolved fluorescence immunoassay (Goldsite diagnostics
kit), colloidal gold immunoassay (Assay Genie POC kit and VivaDia-
COVID-19 IgG-IgM test), ELISA, etc [30]. While a variety of antibody
tests are also available in the market, there is a lot of confusion re-
garding the efficacy of such tests given the high percentage of the
asymptomatic population and the fact that detectable antibodies typi-
cally develop later in the disease. Therefore, diagnostic approach pro-
positions will have to describe their novelty in terms of process, sensi-
tivity, specificity, the strength of risk stratification, and scalability.

We believe that the electrochemical immunosensor technique could
be a potential approach for the POC device application in the detection
of SARS-CoV-2, as shown in (Fig. 6A). There are several recent research
studies about a variety of virus detection using immunosensors
[31–38]. For example, the electrochemical based influenza A virus
HSN1 biosensors are used for the detection of influenza A virus HSN1 to
prevent massive death and to control the transmission from one country
to the other due to their sensitivity, selectivity and economically af-
fordable than the conventional detection method [39]. In another
study, reduced graphene oxide-based electrochemical immunosensor
was developed using EDC-NHS coupling chemistry between the COOH
group of graphene attached on the gold surface of the working ele-
ctrode and NH₂ of antibody specific for H1 of H1N1 influenza and the
reported value of the limit of detection is 0.5 PFU ml⁻¹ [40].

Thus, several studies are still going on based on immunosensor, which
aims to achieve good sensitivity and specificity via different
electrodes surface modifications. In the overall view, the best advantage
of these hypothesized assays is easy handling, no requirement of any
sophisticated instruments, rapid detection efficacy, and cost-effective-
ness than other molecular approaches. Identification of major antigenic
and protective epitopes of target virus particles, is crucial in under-
standing the antibody response while developing these detection stra-
tegies, as antigenic variations make it difficult to generate similar an-
tibodies, thus affecting the development of these diagnostic assays.

5. Aptamer-based detection

Deoxycyribonucleic acid (DNA) is perhaps the most essential of all
biomolecules and has been the foundation for genetic study over the
past few years. However, the use of aptamers, which are small-sized,
single-stranded artificial nucleotides (RNA or DNA), having 10–100
nucleotides, are now gaining enormous importance due to its specific
binding just like in case of antibodies. The selection technique of ap-
tamers is done via the Systematic Evolution of Ligands by Exponential
Enrichment (SELEX) process that binds to a broad range of target
analytes with high affinity and specificity. Aptasensors can measure
analytes in tiny quantities where such small amounts cannot be iden-
tified with most of the other methods being used [41,42]. Furthermore,
the low cost of the aptasensor design compared with other virus diag-
nostic methods should be regarded as one of the economic benefits
[43,44]. Many vital benefits of aptasensors compared to other available
diagnostic techniques include low detection time and a quick detection
method [45,46]. Though aptamers have massive potential as a viable
tool in therapeutics and virus detection, so aptamer-based biosensors
also known as aptasensors, have reported the various methods for the
immobilization of the DNA oligonucleotides sequences and have used
different biomaterials for enhancing the stability and sensitivity of the
electrode [47]. Several electrochemical DNA biosensors have been de-
developed for the detection of various viruses [48–53] and, based on that,
a hypothetical schematic illustration of aptasensor based direct detec-
tion of SARS-CoV-2 using the different platform has been shown in
(Fig. 6B). In a study, DNA influenza virus A H5N1 aptasensor has been
developed, which is a reliable and suitable analytical device with newly
emerged designed for the detection of AIV H5N1 [10]. The active
biological element on the DNA biosensor is specific oligonucleotide
sequences single-stranded DNA (ssDNA) for identifying the viral
genome of complementary ssDNA during the hybridization process.

The standard diagnostic methods implemented for SARS-CoV-2 are
often invasive and costly; these methods often cannot detect the pre-
sence of SARS-CoV-2 virus in tiny amounts. For the identification of
the many viruses with high affinity, aptamers as molecular structures were
used in the aptasensor system. The use of advanced nanostructures and
the functionalized aptamer with certain organic materials will improve
the aptasensor's diagnostic sensitivity and specificity towards SARS-
CoV-2. As the aptasensor specificity and sensitivity are higher, they
make them suitable commercial diagnostic devices for early diagnosis
of SARS-CoV-2. The work is hoped to reduce the effects of interfering
factors in SARS-CoV-2 aptasensors; the more reliable and compact
SARS-CoV-2 diagnostic tools such as commercial aptasensors will also
be developed by incorporating other technologies.

6. CRISPR-based approach

CRISPR is a well-known biotechnological technique for genome
ing editing. Various research groups all over the world have developed a
successful methods for the detection of SARS-CoV-2 using CRISPR.
Zhang et al., developed a protocol for the detection of SARS-CoV-2
using CRISPR-based SHERLOCK (Specific High Sensitivity Enzymatic
Reporter UnLOCKing) technique. Protocol employed the use of Cas13

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for targeting two genes in the virus genome, S gene, Cas13 enzyme and ORF1ab gene. They used synthetic virus RNA fragments as input. The SHERLOCK detection protocol comprises of three steps: 1) isothermal amplification of RNA sample using recombinase polymerase amplification kit, 2) amplified viral RNA is incubated with Cas13 enzyme, guide RNA (gRNA) and reporter. With the help of gRNA, Cas13 enzyme recognizes viral RNA and starts cleaving nearby RNAs, including the reporter. As both ends of the reporter are differently labeled, cleavage creates a unique signature, and 3) cleaved reporter produces distinct band than intact reporter RNA, which can be visualized by eye using the paper dipstick. Using serial dilution of synthetic RNA fragment of the S-gene and ORF1ab gene, Zhang et al., have detected viral RNA in a range between $10^{-100}$ copies/μl [8]. This protocol is developed using synthetic RNA and has not validated using real patient samples. Unlike qPCR, which gives results in hours and requires skilled personnel. SHERLOCK technique is highly sensitive and robust and can give results in less than an hour using dipstick without any elaborate instrumentation.

MammothBiosciences published a white paper on the protocol for rapid detection of SARS-CoV-2 using CRISPR employing DNA endonuclease targeted CRISPR trans reporter (DETECTR). MammothBiosciences reconfigured DETECTR for quick and accurate detection of SARS-CoV-2 using the lateral flow strip. The technique utilizes Cas12 enzyme for targeting three genes, N gene (SARS-CoV-2 specific), E gene (SARS-CoV-2, bat-SARS-like-CoV, and SARS-CoV-2), and P gRNA (sample control). The protocol includes: 1) isothermal amplification of extracted RNA using RT-LAMP 2) Cas12 enzyme incubation to generate RNP complexes, one for each N-gene, E-gene, and P-gRNAs, and reporter probe and 3) visualize result using lateral flow strip. This method can discriminate between SARS-CoV-2 and other coronavirus strains. This assay takes 30 min from the sample to result in the detection limit of 70–300 copies/μl [54].

In the queue of detection of SARS-CoV-2 using CRISPR, Lucia et al., employed Cas12 enzyme for targeting RdRp, ORF1b, and ORF1ab genes, and used WH-human1 sequence as control. They employed synthetic RNA fragments and amplified isothermally using the RPA kit. Amplified fragments were mixed with Cas12, single guide RNA (sgRNA), and reporters. Detection of SARS-CoV-2 was performed using plate reader-based assay and paper strips. ssDNA reporters were labeled with fluorescein amidites (FAM) and biotin for fluorescence and paper assay, respectively. The limit of detection for both assays was 10 copies/μl. The study was carried out using synthetic fragments because of the non-availability of real samples in the nearby regions. The study represents the usefulness of CRISPR-Cas12 technology for the detection of the virus [55].

Besides detection, CRISPR/Cas13 approach can also be used for the treatment of COVID-19. Nguyen et al., proposed that CRISPR/Cas13d can be employed to specifically degrade the SARS-CoV-2 genome, thereby halting its ability to reproduce. For the specificity, they used gRNA containing spacer sequences complementary to ORF1ab and S-gene. With the help of gRNA, Cas13 recognizes the viral genome and chew it up without affecting the human transcriptome. To achieve higher specificity, for the delivery of Cas13d in COVID-19 patients, an adeno-associated virus (AAV) can be used as a vehicle. One AAV can carry up to three gRNAs targeting different genes, thereby increasing the efficiency of treatment. SARS-CoV-2 infects lungs primarily, so to ensure organ-specific delivery, lungs specific AAV serotype can be used. The proposed CRISPR/Cas13d approach is the potentially meteoric approach for the treatment of COVID-19. This approach is under review, and if proven to be effective, it will provide a worldwide weapon.
to fight against RNA virus infection [56].

Abbott et al., developed Prophylactic Antiviral CRISPR in human (PAC-MAN) strategy to define the conserved targetable region of SARS-CoV-2 using CRISPR/Cas13d. They screened and designed a panel of targetable coronavirus RNA. This study revealed that a group of 22 crRNA targets all coronaviruses, whereas a group of 6 crRNA can target 91% of sequenced coronaviruses using CRISPR/Cas13d. Predicted crRNA pool was tested on lung epithelial cell lines using CRISPR because of a lack of access to live SARS-CoV-2 and observed manifold inhibition in viral replication. PAC-MAN strategy could be a powerful tool against COVID-19 [57].

7. Molecularly imprinted polymer (MIP)-based detection

While a variety of bioassays and biosensors have been developed, low-cost, disposable, or reusable biosensors are still needed that can quickly detect and accurately identify the SARS-CoV-2. The use of molecular imprinting technology as elements of biorecognition for the design of MIPs offers a real alternative to antibodies due to their robustness and reproducibility [58]. The molecular imprinting method allows the host components to selectively, sensitively, and rapidly identify and detect the numerous molecules. In the previous years, MIPs based sensors have formed an exciting horizon for surface modification methods by creating specific recognition cavities in the template molecules. They have to turn out to be fascinating, having sensitivity to small structural changes in the biomolecule structure. There are several research reports which are aimed at MIP based approach for medical diagnostics for the detection of various types of viruses such as Influenza virus, Dengue virus, Japanese encephalitis virus, human immunodeficiency virus, Hepatitis virus A and B, Adenovirus, Picorna-virus, etc. [59-66].

For example, a MIPs based nanosensor was developed to detect human papillomavirus derived E7 protein by Cai and co-workers [67]. Their EIS data analysis showed the detection of E7 protein could be as low as sub-pg L⁻¹ levels. Ma et al., have fabricated an electrochemical biosensor based on multi-walled carbon nanotubes modified MIPs for detecting HIV-p24 in human serum samples whose linear range was in the range of 1.0 × 10⁻⁴ to 2.0 ng cm⁻³ and limit of detection (LOD) was found to be 0.083 pg cm⁻³ [68]. Recently, Tancharoen et al., have proposed an electrochemical sensor based on graphene oxide polymers imprinted for the detection of Zika virus. The detection limit of this sensor was then compared to the commercial method and found the observed value to be similar to the reverse transcription [69].

One of the key advantages of MIPs is their robustness, high selectivity, long-term stability, and cost-efficiency, which cannot be accomplished by using fragile biomolecules. MIPs were often used for selective and sensitive viral detection and the distinction of viral subtypes. Thus, it is anticipated that the many advantages of molecular imprinting technology should find its home in the frontlines of possible SARS-CoV-2 detection (Fig. 7).

8. Microarray-based detection

Gene microarray technology can also be used for the diagnosis of SARS-CoV-2. In the microarray approach, there are two main steps: the formation of a specific probe and the production of targeted cDNA fragments. At first, specific probe sequences are immobilized on the glass slide called chip and then-unknown DNA molecules that are present in a sample are cut into fragments by restriction endonucleases. These fragments are attached with fluorescent marker and are allowed to bind with probeesthat are present on the chip. Afterwards the target DNA fragments which have complementary sequences of probes will bind with DNA probe. Hybridization of the fluorescent DNA fragments with a DNA probe on the chip will release fluorescent signals, and analysis is done based on these signals for the identification of DNA fragments. Rong et al., developed a 60-mer oligonucleotide microarray for the detection of SARS coronavirus. Probes were designed, which screen the whole genome of the virus and immobilized on the microarray surface. Samples were taken from the throat swab and gargling fluid of SARS patients, and then RNA was extracted, and cDNA was synthesized by reverse transcription and fragmented through restriction display (RD) technique. Fragmented DNA was labeled with Cy5-universal primer through PCR, and then hybridization was done. There was no signal in negative and blank controls, and results showed that microarray could be used for the detection of SARS coronavirus [70].

As studies find out that there are many point mutations in SARS-CoV genome sequence, Long et al., designed a universal microarray system for detection and genotyping of SARS-CoV by targeting six single nucleotide polymorphisms (SNP) which are allocated through the whole genome. This microarray system was made by combining RT-PCR and ligase detection reaction (LDR). The Zip Codes were attached at the slide surface, and their complimentary Zip Codes were attached with targeted DNA fragments. These Zip Codes sequences were universal as these have no similarity with targeted sequence, human host, or SARS-CoV, so there is no chance of false-positive signal of mismatch hybridization. Twenty samples were tested using this assay, and results were confirmed by DNA sequencing [71]. Lu et al., done screening of specific antigens for the diagnosis of SARS-CoV by using protein microarray. In this study, structural proteins were expressed in E. coli as GST or TRX fusion protein and deposited on the microarray, and testing was done with serum samples from SARS patients. Results show that this GST-N2 fusion protein may prove an essential antigen for serological assay of SARS [72].

Zhang et al., designed a novel method for detection SARS-CoV specific genes by using novel asymmetric multiplex PCR for amplification of DNA with universal primers and labeled PCR products are hybridized with oligonucleotide microarray. Results were analyzed by a fluorescent scanner [73]. Guo et al., developed a novel method: SNP DNA microarray for detection and genotyping of SARS-CoV by targeting 24 SNPs. Sample for testing was taken from 19 SARS-CoV patients, and PCR amplified product was hybridized with the microarray. Results of hybridization show that all samples were detected and genotyped accurately with 100 % accuracy [74]. So, the microarray is a better technique for the detection of the virus with a high mutation rate. Microarray-based diagnostics can be a better approach for detection of SARS-CoV-2 because PCR based methods can detect only a few numbers of genes per experiment, but microarray can detect a higher number of DNA fragments synchronously.

Fig. 7. Hypothetical illustration of MIP based electrochemical sensor for COVID-19 detection. Electrode is fabricated and SARS-CoV-2 could be used as a template to create specific recognition cavities into polymers for their specific detection.
9. Loop-mediated isothermal amplification (LAMP)-based detection

In 2000, Notomi et al., had developed LAMP technique that amplifies DNA quickly with high specificity and sensitivity at a constant temperature [75]. Now, after two decades of its development, LAMP seems a most promising approach to tackle the burden of COVID-19 testing as it is easy to use and does not require specialized machines such as PCR and costly reagents kits. The LAMP test can be done in a single tube at a constant temperature, hence very economical. DNA amplification can be detected by colorimetric, turbidity, or fluorescent-based methods [76]. There are increasing research reports which strongly hint at the possibility of employing the LAMP test for COVID-19 detection. Naama Geva-Zatorsky group from Israel developed a test for COVID-19 detection using Reverse Transcribed Loop-Mediated Isothermal Amplification (RT-LAMP) technique. From the nose and throat swabs, SARS-CoV-2 can be detected by the developed test without RNA purification step [77]. The test was applied to 180 suspected patients, and results were found comparable to standard detection methods for COVID-19.

Another fascinating research report can also be seen on the preprint server (bioRxiv) in which a group from ‘Well Cornell Medicine’ designed a 30-minute colorimetric LAMP test for COVID-19 detection and also developed a shotgun metatranscriptomic profiling platform technology for nose swabs [78]. Simplicity and low cost of the LAMP technique will increase its chance to be chosen as a favored diagnostic tool for COVID-19 detection, and this can be used as a POC device at workplaces, clinics, and entry points.

10. Conclusion

Testing is a valuable tool to control the spread of COVID-19. In our fight with the COVID-19 pandemic, testing is playing a crucial role in public health monitoring. Research reports are evolving rapidly about new diagnostics for COVID-19 detection. Detection technologies based on PCR and antibody are helping us to identify infected people and isolate them for stopping the spread of infection. Currently, detection of COVID-19 is dominated by PCR and antibody-based diagnostics, but alternative technologies such as LAMP, RT-LAMP, CRISPR, etc. are under development and may hit the diagnostic market of COVID-19 soon. Most of the detection technologies use the nasopharyngeal samples for SARS-CoV-2 detection, but oral and blood samples seem more appropriate for upcoming technologies. For sensitivity, reproducibility, and reliability, all COVID-19 testing technologies need to be studied and compared. The next few months will witness the comparison, advantage, and disadvantage of these technologies and how suitable they are for mass screening of healthy and suspected patients. Techniques such as LAMP, which requires minimum resources and reagents may be favoured and increase the testing capabilities of hospitals and clinics. In the coming months, we can witness more comprehensive technologies that will provide more insights for viral and host biology besides just detection. MIP, rolling circle amplification [79] real-time NASBA [80] AI, and smart phone are some examples of future technologies that can be pursued and optimized for the detection of COVID-19. Besides, development of therapeutics and vaccines for the control of COVID-19, the development of new testing technologies is equally important to fight against COVID-19 and lessen the socio-economic harm, we all are facing in current time due to this pandemic.

Declaration of Competing Interest

The authors declare that there are no conflicts of interest.

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