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Conventional PCR assisted single-component assembly of spherical nucleic acids for simple colorimetric detection of SARS-CoV-2

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Abstract
Continuous identification of suspected infectious cases is crucial to control the recent pandemic caused by the novel human coronavirus SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2). Real-time polymerase chain reaction (real-time PCR) technology cannot be implemented easily and in large scale in some communities due to lack of resources and infrastructures. Here, we report a simple colorimetric strategy derived from linker-based single-component assembly of gold nanoparticle-core spherical nucleic acids (AuNP-core SNAs) for visual detection of PCR products of SARS-CoV-2 ribonucleic acid (RNA) template. A palindromic linker is designed based on SARS-CoV-2 specific E gene to program the identical colloidal SNAs into large assemblies along with a distinct red-to-purple color change. The linker acts as a probe of SARS-CoV-2 RNA in conventional PCR reaction. In the presence of the correct template the palindromic linker, which is complementary to a short region within the target amplicon, is cleaved by 5'-exonuclease activity of deoxyribonucleic acid (DNA) polymerase. Cleavage of the palindromic linker during the amplification process inhibits the single-component assembly formation of SNAs. So, positive and negative viral samples produce simply red and purple colors in the post PCR colorimetric test, respectively. Evaluation of the samples obtained from cases with laboratory-confirmed SARS-CoV-2 infection revealed that our assay can rival with real-time PCR method in sensitivity.

1. Introduction
In the late December 2019, an ongoing spread of a viral pneumonia due to the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) was reported in Wuhan, China which quickly erupted into a worldwide pandemic of COVID-19 disease [1–4]. The continuous rise in the number of infected cases and deaths as a result of COVID-19 caused this disease to rapidly become a global public health concern. Extensive screening for COVID-19 has been considered as a crucial strategy in controlling this infectious disease. Unfortunately, due to presenting non-specific influenza-like clinical symptoms or the likelihood of being asymptomatic at onset of illness [2,5,6], it is difficult to early diagnose SARS-CoV-2 infection through clinical symptoms. So, laboratory diagnosis of COVID-19 is the main current issue and challenge for public health systems to prevent the vast spread of the disease.

Computed tomography (CT) imaging is an important tool to diagnose or exclude a disease. Inflammatory lung diseases especially pneumonia cases [7] present abnormalities in lung CT images. Recently, analysis of chest CT images for finding abnormal features has also widely been used for both screening and diagnosis of COVID-19 [5,8–12]. But CT systems cannot specifically diagnose COVID-19 because of large nonspecific overlap of CT imaging features between different viral pneumonias with similar symptoms to COVID-19.

Molecular techniques based on nucleic acid hybridization strategies, in virtue of targeting specific pathogens, are more suitable for accurate diagnosis of COVID-19. The standard molecular method for COVID-19 diagnosis is real-time reverse transcription polymerase chain reaction (rRT-PCR) in which RNA from SARS-CoV-2 is detected. This technique is based on the amplification of complementary DNA (cDNA), synthesized from a conserved region of the viral genome, in a PCR reaction using primer pairs and a dual-labeled fluorophore-quencher probe targeting the cDNA sequence of interest. During the amplification cycles the probe is cleaved, resulting in the fluorescence signal. For molecular techniques to be applied, the complete genetic composition of pathogen (i.e. viral genome sequences) must be ascertained.

SARS-CoV-2 is a single-stranded, positive-sense RNA virus with a
genome of approximately 29.8 kb in size [13] that has been enclosed inside an enveloped virion containing helical nucleocapsid [14]. The coronavirus genomes encode five major open reading frames (ORFs), including a 5′ frameshifted polyprotein (ORF1a/ORF1ab) and four structural genes that encode the four major structural proteins (the spike surface glycoprotein (S), small envelope protein (E), matrix protein (M), and nucleocapsid protein (N)) [13]. It has been proved that in SARS-CoV-2 RNA the same as other RNA viruses, the core residues of viral proteins are highly conserved within structural proteins [15–17].

The early sequencing of the SARS-CoV-2 (Accession number in Genbank: MN908947), allowed for the design of primers and probes needed for the progress of several RT-PCR assays based on targeting different genomic regions, including the RdRp gene (RNA-dependent RNA polymerase, part of the ORF1ab gene), E gene and N gene regions for SARS-CoV-2 specific testing. Among all provided assays, the E gene and RdRp gene assays demonstrated higher detection sensitivity [10, 18–20]. Despite the inherent advantages of real-time PCR, many communities are deficient in required laboratory infrastructures to implement real-time PCR technology. Moreover, the availability of the relevant real-time PCR reagent kits is not balanced with demand for high sample throughput diagnosis to cope with COVID-19 in the time of crisis. Therefore, further investigations for methods to detect SARS-CoV-2 nucleic acid are ongoing.

Nano-technologies rely on the binding between nanoparticles and target molecules to provide a detectable signal for detection of pathogenic species such as SARS-CoV-2 [21–23]. Recently, a combination of plasmonic photothermal (PPT) effect and localized surface plasmon resonance (LSPR) in a dual functional plasmonic biosensor has been used for detection of SARS-CoV-2 nucleic acid [24]. A chip with two-dimensional gold nanoislands functionalized by thiol-complementary DNA receptor of RdRPs, ORF 1ab, or the E gene sequence can detect the selected sequences of SARS-CoV-2 via nucleic acid hybridization. The chip has the capacity of generating local PPT heat that improves the in situ hybridization kinetics of strands for specific nucleic acid detection.

A colorimetric assay for detection of SARS-CoV-2 has also been developed to detect the presence of the virus using the color change of functionalized AuNPs with thiol modified oligonucleotides specific for N-gene (nucleocapsid phosphoprotein) of SARS-CoV-2 [22]. In the presence of the target RNA of SARS-CoV-2, AuNPs modified with thiolated oligonucleotide for nucleocapsid phosphoprotein selectively agglomerate and change in surface plasmon resonance with a red shift in the absorbance spectra happens. This bioassay was totally selective so that no distinct change in absorbance was found with Middle East respiratory syndrome coronavirus (MERS-CoV) RNA. Spherical nucleic acids (SNAs) are a new class of nanostructures, primarily introduced by Mirkin and coworkers [25, 26]. They were made by a dense shell and highly oriented nucleic acids covalently attached to the surface of gold nanoparticles. The binding of DNA to nanoparticles leads to materials which can act as building blocks in assembly processes. SNAs are good diagnostic tools because, through programmable synthesis of localized PPT heat that improves the in situ hybridization kinetics of strands for specific nucleic acid detection.

All designed oligonucleotides, PCR primers and probes were synthesized by Bioneer, Inc. (Korea) with Bio-reversed phase (Bio-RP) purification (except for dual-labeled fluorophore-quencher probe for real-time RT-PCR that was purified by high-performance liquid chromatography (HPLC)). All oligonucleotide sequences are provided in Table 1.

### Table 1

| Oligonucleotide sequences used in this study. |
|-----------------------------------------------|
| DNA name | Sequences and modifications (from 5′ to 3′) |
|---------|--------------------------------------------|
| E gene SNA-DNA⁴ | A*23A24A25A*26A27AAA GTTACACTGATTCCTACCG |
| E gene linker probe⁵ | GGGCAGGAGGATGGCTAGTGTACACAGAGGAGACAGG |
| Forward primer | ACAGGATCGTAAAATGTTAATAGGG |
| Reverse primer | ATATGCGGACGAGTGGACACA |
| E gene real-time probe⁶ | FAM-ACCTAGCCATCCTACTGCCCTGG-23 |

⁴ Phosphorothioate modification is denoted by “*”.
⁵ The underlined bases represent the palindromic sequence.
⁶ FAM: 6-carboxyfluorescein; BBq: blackberry quencher.

### 2. Experimental section

#### 2.1. Instrumentation and apparatus

Ultraviolet-visible (UV–vis) absorption spectra were recorded on a SPECORDER 210 spectrophotometer (Analytik Jena) equipped with a Peltier temperature-controlled cell holder. All spectrophotometric measurements were implemented in a quartz ultra-micro cell cuvette with polytetrafluoroethylene (PTFE) stopper and 1.0 cm path length. Transmission electron microscopy (TEM) images were taken on a Philips EM 208S microscope operating at electron energy of 100 kV. Amplification processes were performed using LightCycler® 96 System from Roche Life Science for real-time RT-PCR and Applied Biosystems™ Veriti™ Dx 96-well from Thermo Fisher Scientific for conventional RT-PCR.

#### 2.2. Chemicals and oligonucleotide sequences

All designed oligonucleotides, PCR primers and probes were synthesized by Bioneer, Inc. (Korea) with Bio-reversed phase (Bio-RP) purification (except for dual-labeled fluorophore-quencher probe for real-time RT-PCR that was purified by high-performance liquid chromatography (HPLC)). All oligonucleotide sequences are provided in Table 1.
Oligonucleotide with phosphorothioate modification of the backbone was used for functionalization of AuNPs to make SNAs. Hydrogen tetrachloroaurate (III) trihydrate, HAuCl₄·3H₂O, 99.99 % and sodium citrate were purchased from Sigma-Aldrich. Sodium chloride, 4-(2-hydroxyethyl) piperazine-1-ethanesulfonate (HEPES), Tris-Borate-EDTA (TBE) buffer (Molecular Biology Grade) and agarose powder for gel electrophoresis were purchased from Merck. Dried oligonucleotides were resuspended in RNase-, and DNase-free water, and ultrapure water was used to synthesize AuNPs. All buffers and other solutions were prepared in ultrapure water.

2.3. Synthesis and characterization of SNA

AuNP cores were synthesized through the standard citrate reduction of HAuCl₄ described elsewhere [33] with slight modifications. Briefly, aqueous solution of 1.0 mM HAuCl₄ was first heated to boil for 20–30 min in a 100 mL round-bottom flask equipped with a condenser. Then, a total of 5.0 mL of 38.8 mM sodium citrate was rapidly added to the boiled HAuCl₄ solution with vigorous stirring. The color of solution changed from pale yellow to deep red in about 1 min. The solution was allowed to reflux for an additional 12 min under vigorous stirring. Then the heating was turned off and the solution was allowed to cool to room temperature under stirring. This colloidal AuNP solution was stored at 4 °C for further use. The functionalization of the synthesized AuNP cores with the desired DNA (SNA-DNA in Table 1) was performed according to the low pH assisted method developed by Liu et al. [34,35] with some modifications. In brief, 500 μL of the already prepared AuNPs was mixed with 15 μL of the stock solution of SNA-DNA (100 μM, resuspended in nuclelease-free water) in an autoclaved microcentrifuge tube by a brief vortexing. After 5 min, 10 μL of 500 mM trisodium citrate–HCl buffer (pH = 3) was added to the AuNP/DNA mixture so that the final concentration of sodium citrate was 10 mM. After 3 min incubation, the concentration of NaCl was brought to 50 mM by adding specific amount of NaCl solution. After another 3 min incubation, 40 μL of HEPES buffer (500 mM, pH 7.6) was added to the tube by a brief vortexing in order to bring the pH value of the AuNP solution back to neutral. To remove the excess amount of non-conjugated DNA, the mixture was centrifuged at 10,000 rpm (revolutions per minute) for 4 min, and the supernatant was removed. The pellet was resuspended in 10 mM HEPES solution. After 3 times repeating this post-conjugation washing, the pellet was resuspended in 300 μL of 10 mM HEPES (in order to adjust the concentration of SNAs around 15 nM for further use). The functionalization of AuNPs with phosphorothioate DNA is facile and rapid so that SNAs can be reproducibly prepared daily or weekly. Characterization of AuNP-core SNAs is available in Supplementary Information (SI) in details.

2.4. Sample collection and RNA extraction

All upper respiratory specimens used in this study were taken from suspected infectious patients who had referred to Farzan Molecular and Pathobiology Laboratory, Hamedan, Iran. According to the declaration of Farzan lab a record of the patients signed their informed consent form for research use is available. Dacron flocked swabs were used to collect samples from upper respiratory tract and the collected samples were kept in the viral transport medium until RNA extraction. Extraction of viral RNA from upper respiratory specimens was performed using the MagNA Pure 96 system (Roche) according to the instructions of manufacturer, and all extracted RNA samples were stored frozen at -80°C until use.

2.5. Amplification process

Reverse transcription and PCR amplification were performed in one step for both the real-time and conventional RT-PCR assays using Add-Probe RT-PCR Kit (Product Code: 74311) and QIAGEN One-Step RT-PCR Kit (Cat. Number: 210210), respectively, according to the manufacturer’s instructions. The same specific primer pair was used for amplifying the E gene in both RT-PCR methods. In a 25-μL total volume of the amplification reaction, the primers were used at an optimized concentration of 0.4 μM, while the linker DNA probe for conventional RT-PCR and the fluorophore-quencher labeled probe for real-time RT-PCR were used at final concentrations of 0.1 and 0.2 μM, respectively. The real-time PCR program with a 20-min reverse transcription step at 50 °C and a 10-min denaturation step at 95 °C to activate the Taq DNA polymerase followed by 45 amplification cycles with a denaturation temperature of 95 °C for 10 s, annealing and extension temperatures of 55 °C for 45 s has been applied. Also, thermal cycling for conventional RT-PCR was performed at 50 °C for 30 min followed by 95 °C for 15 min for reverse transcription and DNA polymerase activation, respectively. Then 45 amplification cycles were run at 94 °C for 15 s, 55 °C for 45 s. Reaction components and optimized concentrations are shown in Table S2.

2.6. Gel electrophoresis

A part of PCR products (7.0 μL) were mixed with 1.0 μL DNA gel loading dye (6X) (Thermo Scientific) and the mixtures were then loaded to different lanes of an agarose minigel (5%) and run at 80 V for 90 min. The 5% agarose minigels were freshly prepared each time before application using 1X TBE buffer, and stained by SYBR Safe dye. TBE buffer (1X) was also used as running buffer. After separation of DNA fragments of the PCR products, the gel images were recorded using a Quantum VilberLourmat gel documentation system. A 50 bp DNA ladder (Invitrogen, Cat. number: 10416014) was used as DNA size marker to determine the band sizes.

2.7. Post-PCR colorimetric assay

The used clinically positive and negative samples had formerly been confirmed by SARS-CoV-2 RNA detection kit (PCR-Fluorescence Probing, EUL 0493-141-00. Da An Gene Co.) according to the manufacturer’s instructions in that ORFlab and N genes were selected as amplification target regions. RNA extraction and conventional RT-PCR amplification (45 cycles) were carried out according to the procedures described in sections 2.4 and 2.5, respectively. Then, the post-PCR colorimetric assay was achieved by mixing 7.5 μL of the PCR product (which contains 2.5 mM MgCl₂ from QIAGEN One Step RT-PCR Kit) with as-prepared SNAs in an eppendorf PCR tube followed by adding NaCl in a total volume of 30 μL (with the final concentration of SNAs, linker probe, MgCl₂, and NaCl 7.5 nM, 25 nM, 0.625 mM, and 380 mM, respectively). After 10 min incubation at 25 °C, the color changes of the solutions were recorded using a digital camera Nikon D7200.

3. Results and discussion

3.1. Design principles of single-component system for SARS-CoV-2 nucleic acid detection

The first step in designing a single-component assembly system based on AuNP-core SNAs is the selection of a linker DNA to play both as oligo-probe for the nucleic acid target and self-assembler of SNA particles. The second step is to design a proper di-block oligonucleotide to form the nucleic acid shell on gold nanoparticles to construct SNAs. The structure of the linker consists of three main parts; a non-palindromic region (region 1, complementary to DNA sequences attached to AuNPs to form the nucleic acid shell on gold nanoparticles to construct SNAs), a palindromic region at the 5′ end (region 2, with self-complementary property), and a non-binding single-nucleotide region, called the flexor (generally adenosine, A) which separates the two regions of the linker and gives flexibility to the linker during assembly process [32,36,37].

To design a proper linker probe, the conserved and specific regions of E gene of SARS-CoV-2 (227-nt E gene sequence) were analyzed to look...
for appropriate palindromic sequences. The palindromic sequences, that could potentially be useful in designing a palindromic linker, are listed in Table S1. We chose the 4-nt 5′-GCGC-3′ sequence (because of faster assembly kinetics) and the 21 nucleotides behind it (towards the 5′ end) in the E gene region for designing a palindromic linker (see Table 1). The selected linker targets the conserved and specific regions of E gene of SARS-CoV-2 RNA and partially overlaps the TaqMan probe in real-time RT-PCR for E gene assay [19,20].

The di-block oligonucleotides comprising SNAs (that we named it SNA-DNA) consist of an attachment moiety of 12-adenines (poly-A) as anchor block and a programmable recognition block that is complementary to the non-palindromic region of the linker. The anchor block and the recognition block are separated by the spacer group, here 4 adenines, which pushes the recognition block away from nanoparticle surface. The anchor block is tandemly modified at the first-eight phosphates by phosphorothioate (PS) where a sulfur atom was substituted for one of non-bridging oxygen atoms on the phosphate backbone. The PS modification has been previously used to immobilize DNA on AuNPs to propose a fast, cost-effective and pre-treatment free method for DNA conjugation [32,35].

3.2. Simulation and characterization of the designed system

After selecting the linker and DNA sequences forming single-component assembly (obtained from 13-nm AuNP core SNAs and the used palindromic linker), we used NUPACK Analysis Algorithms [38–40] to simulate hybridization between the interacting nucleic acid strands can be formed as the most dominant structure (with minimum free energy). Structural evaluation of sequences in NUPACK is done by a polymer graph of a secondary structure. Fig. 1A depicts the polymer graph for the possible secondary structure with minimum free energy (MFE) between the predesigned strands of palindromic linker and SNA-DNAs. The obtained polymer graph demonstrates that the predicted self-complementary assembly structure can be successfully formed via the palindromic base pairing of the linker-hybridized adjacent SNAs.

The self-assembly formation was then investigated by transmission electron microscopy (TEM) and spectrophotometric methods. The mixture of linker/SNAs was deposited on TEM grid, dried and imaged. Remarkably, TEM image demonstrated a flower-like configuration (Fig. 1B) that revealed each particle (i.e. SNA) can be bound to every other SNA with equal possibility. A scheme of the single-component assembly of SNAs is also shown in Fig. 1C for better visualization.

Fig. 1D shows the absorbance spectra vs. time for self-assembly of 13-nm AuNP-core SNAs in the presence of palindromic linker up to 40 min at 25 °C (60 μL of 10 mM HEPES buffer, pH = 7.6, containing 0.45 pmol SNAs, 5 pmol linker and 400 mM of NaCl). The inset is a photograph of the dispersed colloidal SNAs as control and the aggregated SNAs after 10 min of the addition of palindromic linker. Dispersed SNA particles are red, whereas aggregated SNA particles are purple. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).

Fig. 1. Simulative and experimental characterization of the designed single-component assembly system. (A) The polymer graph of the assembly-guided structure simulated by NUPACK Analysis Algorithms (left) and DNA sequences forming single-component assembly (right). (B) TEM image of flower-like structure of single-component assembly. (C) A 3D schematic illustration of flower-like structure of single-component assembly. (D) Time-dependent changes in the UV–vis spectra of 13-nm AuNP-core SNAs in the presence of palindromic linker over 40 min at 25 °C (60 μL of 10 mM HEPES buffer, pH = 7.6, containing 0.45 pmol SNAs, 5 pmol linker and 400 mM of NaCl). The inset is a photograph of the dispersed colloidal SNAs as control and the aggregated SNAs after 10 min of the addition of palindromic linker. Dispersed SNA particles are red, whereas aggregated SNA particles are purple.

(For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).
plasmon band to longer wavelengths. This indicates the formation of larger structures due to self-assembly between linker-hybridized adjacent SNAs through the free sticky ends of the palindromic linker. The kinetics of assembly formation is relatively fast and the color change is visible to naked eyes after 10 min. So, this single-component system can provide sensitive colorimetric responses.

3.3. 5′-exonuclease-based post-PCR assay and workflow for SARS-CoV-2 RNA detection

Exonuclease-based conventional PCR utilizes 5′→3′ exonuclease activity of DNA polymerase as the amplification reaction proceeds through the oligonucleotide probe (here the palindromic linker probe) which cleaves the probe during the extension phase of PCR. Our central idea, captured by the concept of DNA-programmable assembly, is that SNA assembly is inhibited when the linker probe is cleaved in the DNA polymerase amplification process. The used palindromic linker probe is a
short unmodified single-stranded DNA (ssDNA) that is complementary to the sequence of bases in the targeted nucleic acid strand as PCR template. In the presence of target template, the pre-formed duplex of the linker and the target DNA, in the region between the two specific PCR primers of the E gene amplicon, subjected to 5′-exonuclease activity of DNA polymerase [41, 42]. As PCR proceeds through the region where the linker is annealed, the linker DNA is cleaved and removed from its partner during the amplification reaction (Fig. 2A).

The total strategy is graphically represented in Fig. 2B. In general, the upper respiratory tract samples are collected by nasopharyngeal, nasal or throat swabs from the patients. Then, viral RNA is extracted from these specimens. Isolated RNA is reverse transcribed to complementary DNA (cDNA) and subsequently specific regions of cDNA amplified by conventional PCR in the presence of the palindromic linker that is designed as a specific probe for SARS-CoV-2 RNA. In the presence of the target template, cleavage of the linker probe which is formerly annealed to the target template occurs by the 5′-exonuclease activity of the DNA polymerase during the amplification process. Finally, an aliquot of the PCR products is added to the single-component SNA solution (see experimental section). After 5–10 min incubation at room temperature, the colorimetric response can be readily observed by the naked eye. The color of the SNAs keeps red for the positive samples and turns to purple in the case of negative samples.

3.4. Optimization of assay performance

Since our assay provides a post-PCR colorimetric detection, the assay performance is dependent upon both optimum PCR amplification condition and the best colorimetric response associated with the single-component assembly of SNAs. The annealing temperature in PCR amplification process, the ionic strength of the SNA solution and the linker concentration in colorimetric detection are some important factors which must be optimized to get the best performance of the assay. As we targeted the same region in E gene which has been appointed in real-time RT-PCR method for detection of SARS-CoV-2, we used the same target-specific primer pairs [20]. The RNA templates were reverse transcribed using one-step RT-PCR Kit. Then, 45 cycles were performed at different annealing temperatures to amplify the reverse-transcribed DNA templates. Eventually, the PCR products were analyzed using gel electrophoresis (Fig. 3A). The gel electrophoretic analysis of PCR products at the annealing temperatures of 55, 56, 57, and 58 °C, shows a unique-sharp band only in the lane corresponding to the annealing temperature of 55 °C. The details of PCR reaction components are available in Table S2.

In DNA-based nanoparticle assembly, the binding stability of the sticky ends that connects adjacent SNAs together, at a given temperature, mainly depends on the salt concentration. So, the colorimetric
response followed by assembly formation due to self-complementarity of the free palindromic tails is also a function of ionic strength of the medium and the amount of the linker. We observed that with increasing ionic strength and linker concentration, the assembly growth rate also increased. i.e. hybridization of the linker and SNAs became more favorable. Besides the need for stable interconnection of SNAs in the presence of linker for accurate diagnosis, colloidal stability of SNAs alone (with no linker) is another important aspect that also depends on salt concentration [35]. Addition of salt more than the salt tolerance of SNA in solution causes irreversible aggregation of SNAs before analysis. Therefore, the highest salt concentration that provides both colloidal SNAs stability and a fast colorimetric response sufficient for visual detection, in the presence and absence of the correct DNA template, respectively must be determined. On the other hand, the minimum amount of the linker which can sufficiently induce self-assembly of SNAs and observable color change, in the absence of target should also be obtained. This amount helps to get high sensitivity and minimum probability of false negative results in low viral load specimens. As mentioned in section 3.3, the linker is cleaved and removed by the 5′-exonuclease activity of DNA polymerase in conventional PCR amplification process. If high amounts of linker are used, in cases of low viral load samples most of the linker will stay intact and contributes in aggregation of SNA in post-PCR colorimetric detection and leads to false negative results. Therefore, the lowest appropriate amount of linker to get a visual color change in the absence of the correct DNA template and avoid from false negative outcome in low viral load samples must be determined.

To optimize the concentration of NaCl to achieve the above aforementioned condition, the colloidal stability of SNAs at different concentrations of NaCl was first monitored spectrophotometrically. Since the PCR product contains 2.5 mM of MgCl₂ (QIAGEN One Step RT-PCR Kit), the colloidal stability of SNAs was investigated at a given concentration of MgCl₂ (15 µL of 2.5 mM MgCl₂ in 60 µL total volume) while changing the concentration of NaCl up to 460 mM (Fig. 3B). Higher extinction ratio at 700 nm over 520 nm (Ex₇₀₀/Ex₅₂₀ ratio) at greater than 400 mM NaCl, was associated with aggregation of SNAs and irreversible red-to-purple color change. Therefore, in order to ensure the stability of SNAs, the 380 mM was chosen as the optimum concentration of NaCl. Further, in order to optimize the amount of linker, the variations of Ex₇₀₀/Ex₅₂₀ ratio as a function of time were monitored for linker/SNA mixture with various amounts of linker (Fig. 3C). The Ex₇₀₀/Ex₅₂₀ ratios higher than 0.8 leads to a visible color change with the naked eye. So we selected the 1.5 pmol linker (25 nM, total volume 60 µL) to get an easily observable color change in about 10 min. Overall, the optimum performance of our PCR-based colorimetric method was obtained under the following conditions: at the annealing temperature of 55 °C, in the presence of 25 nM of the linker probe for PCR amplification, and 380 mM of NaCl for post-PCR colorimetric detection step in a total volume of 30 µL.

3.5. Verification and validation of the assay

In this work, we used the same specific primers as those commonly used to amplify the E gene sequence of SARS-CoV-2 in real-time RT-PCR. Since the E gene assay has been often recommended for screening SARS-CoV-2, the specificity of the assay has already been established so that no cross-reactivity has been reported with other human coronaviruses [19]. To evaluate the ability of DNA polymerase to degrade the palindromic linker probe (which pre-hybridized with the target template) during amplification of the E gene amplicon, Post-PCR gel electrophoresis was run on 5% agarose gel (Fig. 4A). Whereas, only an intense band of the expected size for the E gene amplicon is observed in the positive control lane, a weak band is observable in the negative control lane with roughly the same size as the band of the linker probe lane. As a proof of the original idea, the colloidal SNAs were added to three PCR tubes containing 25 nM of the palindromic linker, the PCR products of negative and positive control, respectively. As shown in Fig. 4B, a distinct red-to-purple color change is observed in the negative control and linker probe tubes, whereas for the positive control tube, the color of SNAs remains red. These data verify that, in the case of the positive control sample, during the amplification process the annealed linker probe is digested by the 5′-exonuclease activity of DNA polymerase in conventional PCR. However, the linker probe is intact in the negative control sample and the red to purple color changes in the negative control tube indicating the presence of the intact palindromic linkers during the amplification process.

The accuracy of the assay was evaluated in a local clinical lab (Farzan Molecular and Pathobiology Laboratory, Hamedan, Iran) using 9 samples of extracted RNA from patients including 3 positive, 3 low positive, and 3 negative samples that had already been analyzed by the real-time RT-PCR assay. Positive samples are those with positive real-time RT-PCR results targeting amplification of E gene with cycle threshold (Ct) values between 20–25. The low positive samples had been diluted to obtain Ct values between 34–38. All the samples were correctly diagnosed with the proposed assay. To test the assay reproducibility, the PCR product of a positive sample was aliquotted to three individual PCR tubes and frozen at -20 °C. The three replicates were analyzed by freshly prepared SNAs on three different days. All three repetitions gave the same result.

The sensitivity of the method was compared with real-time RT-PCR for the E gene region using serial dilutions (10-fold dilution series with
five dilution steps) of the same extracted RNA samples (Table 2). The real-time RT-PCR and post-PCR colorimetric results have been reported in the form of Ct value and a Boolean answer + or −, respectively. The Ct value is the number of amplification cycles required to detect a real signal from the samples. At this point the target gene exceeds a threshold level and the fluorescence signal emitted during amplification exponentially intersects the threshold line (or background level). The Ct values are inversely proportional to the viral loads so that lower Ct values represent the higher viral RNA copy number. Generally, a Ct value of more than 40 is not clinically defined as a positive result. In contrast, a Ct value less than 40 is interpreted as positive for SARS-CoV-2 RNA. Except for sample 1, the detection performance of real-time RT-PCR in the fifth pipetting step (dilution factor of \(10^5\)) was poor. Whereas, the post-PCR colorimetric method was applicable to all three samples in all five pipetting steps. For further comparison, the amplification curves of real-time RT-PCR, the agarose gel electrophoresis of conventional PCR products, and a photograph of the post-PCR colorimetric detection for five steps of 10-fold dilution series of sample 1 are represented in Fig. 5A–C, respectively. Again the band related to the intact linker is obvious in the negative control lane. As the results imply, it can be claimed that the sensitivity of the post PCR colorimetric method has been increased at least 10 times compared to real-time RT-PCR method.

### 4. Conclusions

Overall, the proposed method is conceptually simple and technically applicable. The simultaneous 5′-exonuclease activity and polymerase activity of the DNA polymerase persuaded us to develop a simple PCR-based platform, which enables practical implementation of the DNA-programmable assembly of SNAs as a colorimetric molecular diagnostic method. This 5′-exonuclease activity allows PCR cycles to digest the assembler linker DNA while amplifying target template DNA. Hence, linker DNA digestion followed by assembly inhibition can be considered as a specific consequence of the presence of the target template DNA. The far-reaching distribution of conventional PCR equipment in most of clinical laboratories makes the conventional PCR an available and

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**Table 2**

| Dilution Factor | Results* | Past-PCR colorimetric method |
|-----------------|----------|------------------------------|
|                 | Real time RT-PCR (Ct value) |                  |
|                 | No. 1     | No. 2     | No. 3     | No. 1    | No. 2 | No. 3 |
| 1               | 21.24     | 23.76     | 23.98     | +        | +    | +     |
| 10              | 23.92     | 26.44     | 27.33     | +        | +    | +     |
| 100             | 26.85     | 30.06     | 31.17     | +        | +    | +     |
| 1000            | 28.31     | 33.07     | 33.97     | +        | +    | +     |
| 10,000          | 33.12     | 36.86     | 37.60     | +        | +    | +     |
| 100,000         | 37.21     | NA        | NA        | +        | +    | +     |

*a* Positive and negative results are indicated by "+" and "−".

*b* NA, Not Applicable.

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**Fig. 5.** Real-time and post PCR analysis of amplification products corresponding to the five-step 10-fold serial dilutions of the positive sample 1. (A) Real-time amplification curves (the dilution factors are shown at the top of each curve). (B) The gel electrophoresis images and, (C) Photographs showing colorimetric response (pure linker probe was implemented alongside the serial dilutions).
affordable platform for molecular diagnosis of viral pathogens. Therefore, our PCR-based colorimetric method can partly contribute to meet the demand for wide-scale screening for suspected infected patients. In this paper, our PCR-based colorimetric method can partly contribute to meet the demand for wide-scale screening for suspected infected patients. In this paper, our PCR-based colorimetric method can partly contribute to meet the demand for wide-scale screening for suspected infected patients.
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