Temperature responsive smart polymer for enabling affinity enrichment of current coronavirus (SARS-CoV-2) to improve its diagnostic sensitivity

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Abstract
The current commercially available SARS-CoV-2 diagnostic approaches including nucleic acid molecular assaying using polymerase chain reaction (PCR) have many limitations and drawbacks. SARS-CoV-2 diagnostic strategies were reported to have a high false-negative rate and low sensitivity due to low viral antibodies or antigenic load in the specimens, which is why even PCR test is recommended to be repeated to overcome this problem. Thus, in anticipation of COVID-19 current wave and the upcoming waves, we should have an accurate and rapid diagnostic tool to control this pandemic. In this study, we developed a novel preanalytical strategy to be used for SARS-CoV-2 specimen enrichment to avoid misdiagnosis. This method depends on the immuno-affinity trapping of the viral target followed by in situ thermal precipitation and enrichment. We designed, synthesized, and characterized a thermal-responsive polymer poly (N-isopropylacrylamide-co-2-hydroxyisopropylacrylamide-co-strained alkyne isopropylacrylamide) followed by decoration with SARS-CoV-2 antibody. Different investigations approved the successful synthesis of the polymeric antibody conjugate. This conjugate was shown to enrich recombinant SARS-CoV-2 nucleocapsid protein samples to about 6 folds. This developed system succeeded in avoiding the misdiagnosis of low viral load specimens using the lateral flow immunoassay test. The strength of this work is that, to the best of our knowledge, this report may be the first to functionalize SARS-CoV-2 antibody to a thermo-responsive polymer for increasing its screening sensitivity during the current pandemic.

1. Introduction
In anticipation of SARS-CoV-2 current wave and the upcoming waves, we should have an accurate and rapid diagnostic tool that will be critical in controlling the pandemic, which is expected to cause severe attacks, especially by changing whether temperature [1]. The current coronavirus disease 2019 (COVID-19) commercially available diagnostic approaches include two categories, the first one depends on viral RNA molecular assaying using polymerase chain reaction (PCR) based techniques and the second is the serological and immunological assays that detect antibodies or antigenic proteins in patient’s specimens after viral exposure, as discussed recently by our group [2]. Unfortunately, most of the currently available diagnostic approaches have many limitations and drawbacks, even SARS-CoV-2 nucleic acid molecular assaying using polymerase chain reaction (PCR) based techniques that are considered as the gold standard methods for assaying the virus was reported to have a high false-negative rate and low sensitivity [3,4]. The current SARS-CoV-2 RT-PCR positive rate of detection is only 30%-50% and the viral RNA detectability is very limited between days 15 to 39 after the onset of infection [5,6].

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Different factors contribute to SARS-CoV-2 RT-PCR false-negative results, but the sampling techniques and sample type are the common factors where nasopharyngeal and oropharyngeal (NP/OP) specimens are the common samples used in this assay despite the fluctuation in viral load according to the stage of infection where SARS-CoV-2 colonize in the lower respiratory tract [7–10]. Most COVID-19 patients cannot be diagnosed before premonir because of the low viral load in the specimens [9,11–13].

SARS-CoV-2 viral load change is similar to influenza, but completely different from the change in SARS and MERS, comparable to chest CT scan, SARS-CoV-2 viral load peaked approximately at day 10 after the onset of infection [14–16]. Therefore, the low number of viral copies in the specimen increases the rate of false-negative PCR assay and virus misdiagnosis, that is why even the PCR test is recommended to be repeated several times to overcome this problem.

On the other hand, the presence of SARS-CoV-2 antibodies and antigenic proteins in plasma was reported to be <40% among the individuals after the first week of infection onset [17]. Viral antibody immune response may not be enhanced in some patients until the 4th week after the onset of infection, especially in asymptomatic patients, and that means SARS-CoV-2 negative results should not affect patient management and infection control decisions especially, these misdiagnosed peoples are active carriers of the virus and failure to quarantine them would be a great setback in SARS-CoV-2 transmission [18–21].

Thus, it is now clear that our great challenge regarding the current pandemic diagnosis is how to avoid the false-negative diagnosis that may be caused by low viral antibodies or antigenic loads in the analyzed specimens regardless of the variations of the currently available diagnosis techniques, specimen type, sampling technique, and infection stage. In other words, we are in big need of a novel preanalytical strategy to be used for specimen enrichment to avoid misdiagnosis.

Immuno-affinity trapping strategies that mainly depend on antigen-antibody interaction have been confirmed as effective methods in concentrating trace biomarkers in various body fluids and biological samples [22]. Stimuli-responsive polymers decorated with specific antibodies have been used for the immune perception of different biomolecules in different biomedical applications, especially immunoassaying, and others depending on the reversible aggregation after induction of temperature, pH, light, etc. [23–25]. Our group possesses many achievements in this trend [26–31].

In this study, we designed, synthesized, and characterized a temperature-responsive polymer, poly(N-isopropylacrylamide-co-2-hydroxysopropylacrylamide-co-strained alkyl isopropylacrylamide) (P[NIPAAm-co-HPAAm-co-SAKIPAAm]) followed by decoration using anti-SARS-CoV-2 nucleocapsid antibody for enabling affinity enrichment of recombinant SARS-CoV-2 nucleocapsid protein through in situ precipitation process as an initial model to be used as a pre-analytical technique to increase targets concentrations in SARS-CoV-2 assays to overcome misdiagnosis and improving its diagnostic sensitivity as shown in Fig. 1.

2. Materials and methods

2.1. Materials

N-isopropylacrylamide (NIPAAm, Wako Pure Chemical, 97%) was recrystallized from hexane and dried under vacuum before use. chloroform (Wako Pure Chemical, 99.9%), D,L-2-amino-1-propanol (Tokyo Kasei, 98.0%), triethyamine (Wako Pure Chemical), acryloyl chloride (Tokyo Kasei, 95.0%), 2-propanol (Wako Pure Chemical, 99.7%), ethyl acetate (Kishida chemical, 99.5%), cyanomethyl dodecyl trithiocarbonate (CDT, Aldrich), 2,2′-azobis(isobutyronitrile) (AIBN, Wako Pure Chemical), tetrahydrofuran (THF, Kanto chemical, 99.0%), diethyl ether (Wako Pure Chemical, 99.5%), deuterium oxide containing 0.05 v/v% sodium 3-(trimethylsilyl)-1-propanesulfonate-d6 (TMSP) (D2O, Wako Pure Chemical, 99.9%), dimethyl sulfoxide-d6 containing 0.05 v/v% tetramethylsilane (TMS) (DMSO d6, Wako Pure Chemical, 99.9%), N,N-dimethylformamide (DMF, Wako Pure Chemical, 99.7%), dibenzyclooctyloctyacetic acid (DIBAC, Click Chemistry Tools, 95.0%), dichloromethane (DCM, Wako Pure Chemical, 99.0%), goat anti-human IgG antibody (abcam), SARS-CoV-2 nucleocapsid humanized antibodies, >95% (MBS355887, Clone #5B, MyBioSource, Inc, San Diego, USA), Recombinant SARS-CoV-2 nucleocapsid Protein, >95% (MBS355892, MyBioSource, Inc, San Diego, USA), Pierce® modified lowry protein assay kit (thermo fisher scientific), 4-dimethylamino pyridine (DMAP, Wako Pure Chemical), N,N′-dicyclohexylcarbodi mide (DCC, Tokyo Kasei, 98.0%), ethanol (Wako Pure Chemical, 99.5%), Dulbecco’s phosphate buffered saline (PBS, Aldrich), azido-ethylene glycol (EG4)-NHS ester (Tokyo Kasei), dimethyl sulfoxide (DMSO, Wako Pure Chemical, 99.0%), copper(II) sulfate, 5-hydrate (Aldrich), DL-2-aminoobutyric acid (Tokyo Kasei, 99.0%), fluorescamin (Tokyo Kasei) - methanol (99.8% - Wako Pure Chemical), 10 × tris/glycine/SDS buffer (BIO-RAD), Coomassie brilliant blue R-250 (CBB, BIO-RAD), Laemmlli sample buffer (BIO-RAD), 2-mercaptoethanol (Wako Pure Chemical, 99%), Precision plus protein unstained standards (BIO-RAD).

2.2. Synthesis and characterization of HIPAAm monomers

HIPAAm was synthesized according to the previous report [32], where D,L-2-amino-1-propanol (0.15 mol), and triethyamine (0.15 mol) were dissolved in anhydrous chloroform solvent followed by stirring at 5 °C for 20 min then acryloyl chloride (0.15 mol) was added to the prepared solution very slowly, followed by stirring at 5 °C for 2 h. The obtained solvent was evaporated using the Rota evaporator followed by re-dissolving in 2-propanol and keeping at −20 °C for more than 24 h. Finally, filtration was performed for salts removal followed by concentration and purification using column chromatography. Synthesis of HIPAAm monomer was confirmed using thin-layer chromatography (TLC) followed by 1H NMR (Solvent D2O).

2.3. Synthesis of P(NIPAAm-co-HIPAAm)

P(NIPAAm-co-HIPAAm) was synthesized by Reversible Addition-Fragmentation Chain Transfer Polymerization (RAFT polymerization) of HIPAAm hydroxyl derivative with NIPAAm (temperature-responsive polymer) as shown in Scheme 1(I). Reaction conditions were as the following (NIPAAm 1.89 g, HIPAAm 0.11 g, AIBN 1.31 mg, CDT 12.7 mg, ethanol 17.6 ml) followed by stirring at 20 °C for 20 h, evaporation, and vacuum drying.

2.4. Synthesis of P(NIPAAm-co-HIPAAm-co-SAKIPAAm)

Dibenzyclooctyloctyacetic (DIBAC) acid that contains an alkylene group was introduced into HIPAAm hydroxyl group by dehydration, condensation to obtain our clickable responsive polymer P(NIPAAm-co-HIPAAm-co-SAKIPAAm) as shown in Scheme 1(II). Reaction conditions were as the following (DCM 30 ml, DBCO acid 35.8 mg, DMAP 12 mg, P(NIPAAm-co-HIPAAm) 100 mg, DCCD 20 mg) followed by stirring overnight, evaporation, and vacuum drying.
2.5. Characterization of the polymeric synthesis

Both P(NIPAAm-co-HIPAAm) & P(NIPAAm-co-HIPAAm-co-SAKIPAAm) were confirmed by $^1$H NMR (Solvent:D$_2$O, DMSO), Gel permeation chromatography (GPC) (Solvent:DMF, Standard: PS), Lower critical solution temperature (LCST) (Solvent: PBS, pH: 7.4, conc.: 2.0 mg/mL, temp.: 10–40°C).

2.6. Introduction of azido group to the antibody

Azido-PEG$_4$-NHS ester powder was dissolved in DMSO (10 mg/ml) and different concentrations of Azido-PEG$_4$-NHS ester were prepared using carbonate buffer (pH 8.6). Azido-PEG$_4$-NHS ester was added to the antibody with different feeding ratios followed by rotation at 4°C for 5 h as shown in Scheme 1(III).

2.7. Evaluation of azido group introduction to the antibody

After conjugation of the azido group to the antibody, we measured the fluorescence of the antibody samples using Infinite® 200 PRO plate reader as following: Azido modified antibody samples were diluted to 0.5 mg/ml and then applied to a 96 microwell plate beside different concentrations of D,L-2-Aminobutyric acid that were used as standards (50 µL/well then 5 µL of fluorescamine (50 mg/ml) was added to each well followed by incubation in dark for 15 min and fluorescence investigation (395 nm/495 nm). All samples were applied as triplicates.

2.8. Preparation of SARS-CoV-2 antibody-temperature-responsive polymer conjugate via click reaction

The azido-modified antibody was conjugated to the alkyne group modified temperature-responsive polymer using click chemistry as following: 75 µL of P(NIPAAm-co-HIPAAm-co-SAKIPAAm) was mixed with 50 µL of azido-modified antibody (2.5 mg/ml in PBS) and stirred at 4°C for 12 h as shown in Scheme 1(IV). The polymer concentration was adjusted for the following antibody: polymer ratios – 1:1, 1:2, 1:4, 1:8, 1:15, and 1:30. The succeeded conjugates were characterized by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE) compared with the antibody and polymer.

2.9. Evaluation of SARS-CoV-2 antibody-temperature-responsive polymer enrichment efficacy

The previously prepared SARS-CoV-2 antibody-polymer conjugate (1:1) ratio enrichment efficacy with free polymer was evaluated by thermal precipitation. Purified SARS-CoV-2 -polymer conjugates (1.3 mg/mL, 100 µL in PBS with 15 equivalents of free polymer) were added to microtubes and centrifuged (13000×g) for 5 min at 37°C. Then the supernatant (90 µL) and precipitate (10 µL) were collected, and the absorbance at 280 nm was measured by UV-vis. The molar ratios of SARS-CoV-2 antibodies in the liquid phase and the solid phase were estimated, and the enrichment ratios were calculated.

2.10. Evaluation of SARS-CoV-2 antibody-temperature-responsive polymer antigenic enrichment efficacy

Recombinant SARS-CoV-2 nucleocapsid protein enrichment capacity was evaluated against different concentrations of purified SARS-CoV-2 antibody polymer conjugates (1.3, 0.7, 0.35, 0.17, 0.08, 0 mg/mL, 50 µL in PBS) with 15 equivalent of the free polymer, where antibody-polymer conjugate (different concentrations, 50 µL in PBS) were mixed with the SARS-CoV-2 recombinant protein (1.0 mg/mL, 50 µL in PBS), and incubated for 1 h followed by addition of the free polymer to the solution and centrifugation in microtubes at 37°C, 13000×g for 5 min. The supernatant (90 µL) and precipitate (10 µL) were collected, and the SARS-CoV-2 recombinant protein in the supernatant was measured using the modified Lowery protein assay kit according to the manufacturer instructions and finally, the enrichment ratios were calculated.

2.11. Evaluation of SARS-CoV-2 antibody-temperature-responsive polymer antigenic enrichment efficacy using lateral flow immunoassay strip

Different concentrations of purified SARS-CoV-2 recombinant protein (1.04 × 10^{-15}–20.83 × 10^{-15} mol/mL, 20 µL in PBS) was tested using lateral flow immunoassay strips (that were obtained from Prof. Gamal Shiha, Egyptian Liver Research Institute and Hospital, Mansoura, Egypt) for the determination of the detection limit of SARS-CoV-2 lateral flow-immunoassay strip (LFIA) then
the samples that were detected negative using LFIA were retested after using our devolved system with free polymers as following: purified SARS-CoV-2 recombinant protein (1.04 × 10⁻¹⁵ and 2.08 × 10⁻¹⁵ mol/ml,100 μL in PBS were mixed with the purified SARS-CoV-2 antibody-polymer conjugate, and incubated for 1 h followed by addition of 15 equivalent of the free polymer and centrifugation in microtubes at 37 °C, 13000 × g for 5 min. The supernatant (180 μL) and precipitate (20 μL) were collected, the enriched part was assayed using LFIA.

2.12. Validation of the designed material strategy and polymeric conjugation in real settings

(P(NIPAAm-co-HIPAAm-co-SAKIPAAm)) and azido modified antibodies were mixed and allowed to react according to previous conditions using different real biological samples compared with PBS to evaluate the limitations that may be happened due to sample complexity and heterogeneity. PBS, nasopharyngeal samples, oropharyngeal samples, and mid-stream urine were used as the
solvents. Nasopharyngeal, oropharyngeal, and urine samples were obtained from a healthy volunteer, samples were collected in sterile containers and used freshly in our experiments. Handling of these samples was performed according to the International Ethical Guidelines for Biomedical Research Involving Human Subjects (CIOMS/WHO, 1993). Finally, conjugates conducted in different samples were characterized using SDS-PAGE compared to the antibody and polymer.

2.13. Statistical analysis

Data were analyzed using the Statistical Package of Social Science (SPSS) program for Windows (Standard version 21). The descriptive statistics were presented as mean ± SD (standard deviation) for parametric data. ANOVA test was used to compare more than 2 means, and paired t-test was used to compare paired data. The threshold of significance was fixed at the 5% level (P-value). Results were considered significant when the probability of error was <5% (P < 0.05).

3. Results

3.1. Confirmation of HIPAAm synthesis

The purification of HIPAAm monomer was confirmed by the TLC analysis Supplementary Information A (Fig. S1) that showed that HIPAAm was successfully prepared with two main secondary products. Column chromatography was used for HIPAAm purification as shown also in Fig. S1, and then finally, confirmation of HIPAAm synthesis was done using 1H NMR results (Fig. S2).

3.2. Confirmation and characterization of P(NIPAAm-co-HIPAAm) & P (NIPAAm-co-HIPAAm-co-SAKIPAAm)

1H NMR analysis showed and confirmed the successful polymerization of HIPAAm with NIPAAm with the ratio 3.6: 96.4, where PNIPAAm LCST that is well known to be around 32 °C was increased after introducing HIPAAm hydroxy group to be 37.4 °C and the successful conjugation of the strained alkyne, SAK group to form our temperature-responsive polymer P(NIPAAm-co-HIPAAm-co-SAKIPAAm) with ratio 96.4:1.2:2.4. Moreover, regarding the molecular weight, GPC analysis showed that the molecular weight of the synthesized polymer was shifted from 1.904 × 10^4 to 2.014 × 10^4 (g/mol) due to the insertion of the strained alkyne, SAK group and the LCST was shifted from 37.4 to 30.1 °C as shown in Figs. S3–S5, Fig. 2 & Table 1.

3.3. Synthesis of azido-Anti- SARS-CoV-2 antibody

Introduction of Azido-groups to Anti- SARS-CoV-2 antibody was performed by binding of azido-[EG]4-NHS to the antibody lysine residues. Azido-[EG]4-NHS confirmation and quantification were done using the fluorescamine reduction method where more conjugation results in fewer available lysine residues and amine groups and vice versa. Increasing the azido-[EG]4-NHS feeding resulted in the gradual increase of azido group conjugation as follows (0, 6.7, 10.7, 20.9) and gradual decreasing of amine group conjugation as follows (21.1, 14.7, 10.7, 0.2) p-value < 0.001 as shown in (Fig. 3).

3.4. Confirmation of the synthesized Anti- SARS-CoV-2 polymer conjugate prepared via click reaction

Sodium dodecyl sulfate gel electrophoresis (SDS-PAGE) gel images of different concentrations of polymer with a constant concentration of anti-SARS-CoV-2 antibody. Different anti-SARS-CoV-2 antibody: polymer ratios – 1:1, 1:2, 1:4, 1:8, 1:15, and 1:30 (lanes 2 to 7) were investigated as shown in Fig. 4, increasing the amount of polymer in the reaction solution caused more conjugation to the antibody and finally, more band broadening and increasing in the conjugate molecular weight. Compared to lane 8 that showed no band where it contained only free polymer and (lane 9:10) that showed the denatured light and heavy chains of both free SARS-CoV-2 antibody and azido SARS-CoV-2 antibody. These results indicate successful anti-SARS-CoV-2 conjugated polymer synthesis.

3.5. SARS-CoV-2 antibody-temperature-responsive polymer enrichment efficacy evaluation

SARS-CoV-2 antibody-temperature-responsive polymer supported with free polymer showed a highly significant enrichment capacity after applying thermal precipitation compared to the control one without applying thermal precipitation, p-value < 0.001 as shown in Fig. 5 where no. of enrichment folds was increased to reach around 71-fold after thermal induction.

3.6. Antigen enrichment using SARS-CoV-2 antibody-temperature-responsive polymer

As shown in Fig. 6, by increasing the concentration of the purified SARS-CoV-2 antibody polymer conjugate, the number of recombinant SARS-CoV-2 nucleocapsid protein folds were shifted from 0 to reach to about 6 folds of the original antigenic concentration with the highest conjugate concentration (1.3 mg/mL) p-value < 0.05 that indicated the capability of our developed system to enrich low viral load samples.

3.7. Evaluation of antigen enrichment using lateral flow immunoassay strip

As shown in Fig. 7A, the minimum detection limit of recombinant SARS-CoV-2 nucleocapsid protein (48 kDa) using LFIA was around 1.08 × 10^{-15} mol/mL (a5) and 2.08 × 10^{-15} mol/mL (a4) showed a weak faint band at the test zone while concentrations 4.17 × 10^{-15} to 20.83 × 10^{-15} mol/mL showed strong positive bands as shown in Fig. 7a1, a2, a3 respectively. However, both false-negative concentrations showed strong positive bands after enrichment using SARS-CoV-2 antibody polymer conjugate as shown in Fig. 7B

3.8. The designed material strategy can be proceeded regardless of body fluid heterogeneity

As shown in Fig. 8, Sodium dodecyl sulfate gel electrophoresis (SDS-PAGE) gel image for the designed polymer conjugation in different body fluids including nasopharyngeal, oropharyngeal, and urine samples compared to PBS (lanes 5–12) and compared with antibody alone, polymer conjugates light and heavy chains were shifted to higher molecular weights, with a marked band broadening and smeared migration patterns in the loading well. Compared to lane 4 that showed no band where it contained only free polymer and (lane 2:3) that showed the denatured light and heavy chains of both free antibody and azido antibody. These results indicate successful polymeric conjugation in different body fluids including the upper respiratory tract samples that are commonly used for SARS-CoV-2 detection regardless of the heterogeneity and complexity of these samples.

4. Discussion

Most of SARS-CoV-2 commercially available diagnostic methods including PCR, ELISA, and LFIA based techniques have many
limitations, especially high false-negative rates and poor sensitivity [2–4]. Low viral load in the tested specimens is the main cause of this misdiagnosis [8,9]. Especially, nasopharyngeal and oropharyngeal (NP/OP) specimens are the common samples used for SARS-CoV-2 detection despite their problems [10]. Our findings were in agreement with these previous reports Supplementary Information(B).

On the other hand, the special nature of the current pandemic affects this situation where SARS-CoV-2 viral load peaked approximately at day 10 after the onset of infection [14–16]. SARS-CoV-2 antibodies and antigenic proteins in blood plasma were reported to be<40% among individuals after the first week of infection onset [17]. Viral antibody immune response may not be enhanced in some patients until the 4th week after the onset of...
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Poly NIPAAm (PNIPAAm), is a temperature-responsive polymer that was selected in our study for its water solubility below its LCST of 32 °C where the PNIPAAm hydrophilic amide conjugates with water molecules by hydrogen-bonding and water molecules surrounding the hydrophobic isopropyl group performs hydrophobic hydration by hydrogen bonds causing the complete water solubility [33]. On the other hand, it aggregates above the LCST 32 °C where the hydrophobic hydration turns to an unstable state by increasing the temperature above 32 °C, causing PNIPAAm dehydration and aggregation due to hydrophobic interaction [34].

Therefore, depending on the previous, we can easily control the solubility state of PNIPAAm in water by thermal induction. Moreover, decoration of PNIPAAm with specific antibodies can enable immuno-affinity trapping and finally enrichment of different antigenic targets by applying thermal induction.

In our study, we used click chemistry for enabling polymer and SARS-CoV-2 antibody conjugation. Therefore, we labelled anti-SARS-CoV-2 nucleocapsid antibody with azido groups and conjugated the thermo-responsive polymer with a cyclic strained alkyne group (SAK group) for enabling the Huisgen cycloaddition click reaction [35] between the antibody azido group and SAK group. Click reaction forms a triazole ring by a 1,3-dipolar cycloaddition of the azido group and the alkyne group [36]. This reaction is highly selective and specific and can proceed regardless of the surrounding reaction environment including impurities, temperature, and reaction solution [37]. These properties explain why we preferred the click chemistry for the polymeric antibody conjugation.

In this study, P(NIPAm-co-HIPAm) was prepared using RAFT polymerization to enable the introduction of the strained alkyne group into NIPAAm. The synthesis was confirmed using 1H NMR, also, LCST measurements confirmed the polymeric synthesis where the PNIPAAm LCST that is well known to be around 32 °C was increased after introducing HIPAAm hydroxy group to be 37.4 °C and decreased by introducing the hydrophobic SAKIPAAm strained alkyne group to be 30.1 °C in addition to GPC measurements that confirmed the molecular weight change. To the best of our knowledge, this may be the first report that introduces the strained alkyne group into NIPAAm copolymer, but these findings were agreed with our previous report where we introduced carboxy isopropyl acrylamide into the NIPAAm copolymer and mentioned the relative LCST elevation [38] also, our findings were parallel with Maeda et al., who introduced HIPAAm into the NIPAAm copolymer [39]. SARS-CoV-2 antibody conjugation with azido groups was achieved by the addition of azido-(EG)₄-NHS to the antibody lysine residues that were confirmed by the fluorescamine reduction method where increasing azido-(EG)₄-NHS feeding during the reaction showed a higher azido conjugation to the antibody and fewer amine residues, these results were in agreement with Byeong results who labelled the anti-Her2 IgG to azido groups previously [40]. Our SDS-PAGE gel images showed different polymeric concentrations after conjugation with a constant concentration of SARS-CoV-2 antibody. Increasing the amount of polymer in the reaction solution caused more conjugation to the antibody and finally, more band broadening and increasing in the conjugate molecular weight, which confirmed the successful conjugate synthesis. Finally, we decided to use the antibody to polymer ratio 1:1 to avoid any kind of antibody denaturation due to over polymeric conjugation and to keep the antibody affinity as much as we can in combination with the free polymer to enhance conjugate sedimentation, these data were inconsistent with Huisgen click reaction findings [33].

SARS-CoV-2 antibody-temperature-responsive polymer supported with free polymers showed a high significant recovery after applying thermal precipitation that enabled recombinant SARS-CoV-2 nucleocapsid protein enrichment to about 6 folds of its original concentration that indicated the capability of our developed system to enrich low viral load samples, these data were parallel

infection, especially in asymptomatic patients [18-21]. Thus, our great challenge during this third wave and the upcoming waves is how to reduce the misdiagnosis rates of COVID-19 patients to decrease viral transmission.
concentrations 1.04 and 2.08 would have a great setback in SARS-CoV-2 transmission. The two methods with different respiratory tract samples and this problem discussed the same obstacle especially, during using LFIA diagnostic these results were inconsistent with previous reports [43–46] that CoV-2 samples, especially low viral load specimens happened, and pandemic. According to Fig. 7A & B, misdiagnosis of some SARS-CoV-2 antibody-temperature-responsive polymer efficacies, Fig. A Different concentrations of recombinant SARS-CoV-2 nucleocapsid protein was tested using lateral flow immunoassay, a1: 20.83 × 10^{-15}, a2: 10.42 × 10^{-15}, a3: 4.17 × 10^{-15}, a4: 2.08 × 10^{-15} and a5: 1.04 × 10^{-15} mol/mL. Fig. B Comparing samples before and after enrichment using SARS-CoV-2 antibody-temperature-responsive polymer, b1: 1.04 × 10^{-15} mol/mL without enrichment, b2: 1.04 × 10^{-15} mol/mL after enrichment, b3: 2.08 × 10^{-15} mol/mL without enrichment, b4: 2.08 × 10^{-15} mol/mL after enrichment.

Our system succeeded in antigenic enrichment and avoiding misdiagnosis of low concentration specimens. The two concentrations 1.04 and 2.08 × 10^{-15} mol/mL were detected as negative samples, then detected as positive after our polymeric system treatment. Body fluids are complexed and heterogeneous solutions that contain a massive content of different biomolecules and impurities that may affect polymeric conjugation negatively [47]. To overcome this challenge without any adverse effects on the polymeric conjugation and affinity, we conducted our experiments using click chemistry that is considered highly selective and specific and can proceed regardless of the reaction environment [37]. This concept was clear in our findings where the results indicated successful polymeric conjugation in different body fluids including the upper respiratory tract samples that are commonly used for SARS-CoV-2 detection. Polymeric conjugation was succeeded in urine, nasopharyngeal, and oropharyngeal specimens regardless of the heterogeneity and complexity of these samples. This report may be the first to conduct polymeric conjugation for diagnosis purposes in real body fluids, especially upper respiratory tract samples that commonly used for SARS-CoV-2 detection.

The limitation of this work may be that we need to functionalize our polymeric strategy using not only anti-SARS-CoV-2 nucleocapsid antibody but also different antibodies to target different viral epitopes, especially some antigenic targets are hidden or highly glycosylated in native form. Moreover, we need more validation for these different SARS-CoV-2 antigenic targets to be used in many diagnostic approaches, especially PCR.

5. Conclusion

In our study, we developed a portable, fast, easy, and sensitive strategy based on the concepts of immune-affinity and responsive smart polymeric materials. This strategy was developed to be used as a preanalytical step for SARS-CoV-2 specimen enrichment to avoid the misdiagnosis that usually happens during SARS-CoV-2 diagnosis due to the low viral load in the tested samples. Our developed system succeeded in antigenic enrichment where it enriched recombinant SARS-CoV-2 nucleocapsid protein samples to about 6 folds in addition to avoiding misdiagnosis of low viral load specimens that were detected as negative samples, then detected positive after our polymeric system treatment. The strength of this work is that to the best of our knowledge, this report may be the first to try to functionalize SARS-CoV-2 antibodies to a thermo-responsive polymer for utilization of increasing the screening sensitivity during the current pandemic.
akab: Investigation, Methodology, Formal analysis, Writing - review & editing. A.A. Hassan: Formal analysis, Writing - review & editing.
G. Shiha: Formal analysis, Writing - review & editing. M. Ebara: Supervision, Resources, Funding acquisition, Project administration, Conceptualization, Data curation, Formal analysis, Validation, Visualization, Writing - original draft, Writing - review & editing.

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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Appendix A. Supplementary data

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