1	Development of RNA-based assay for rapid detection of SARS-CoV-2 in clinical samples				
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24					

26 Abstract:

27 The ongoing spread of pandemic coronavirus disease (COVID-19) is caused by Severe Acute 28 Respiratory Syndrome coronavirus 2 (SARS-CoV-2). In the lack of specific drugs or vaccines for 29 SARS-CoV-2, it demands rapid diagnosis and management for controlling the outbreak in the 30 community. Here we report the development of the first rapid-colorimetric assay capable of detecting SARS-CoV-2 in the human nasopharyngeal RNA sample in less than 30 minutes. We 31 32 utilized a nanomaterial-based optical sensing platform to detect RNA-dependent RNA polymerase (RdRp) gene of SARS-CoV-2, where the formation of oligo probe-target hybrid led 33 34 to salt-induced aggregation and changes in gold-colloid color from pink to blue in visible range. Accordingly, we found a change in colloid color from pink to blue in assay containing 35 nasopharyngeal RNA sample from the subject with clinically diagnosed COVID-19 (n=7). The 36 37 colloid retained pink color when the test includes samples from COVID-19 negative subjects 38 (n=6) or human papillomavirus (HPV) infected women (n=2). The optimized method has detection limit as little as 0.5 ng of SARS-CoV-2 RNA. Overall, the developed assay rapidly 39 detects SARS-CoV-2 RNA in clinical samples in a cost-effective manner and would be useful in 40 41 pandemic management by facilitating mass screening.

42

43 Key words

44 COVID-19, diagnosis, Ribonucleic acid, Coronavirus disease, colorimetric test

46 Introduction

Coronavirus disease is rapidly spreading across the world and raising severe global health 47 concerns. In December 2019, China reported the first disease case in its Hubei Province. Based 48 49 on the phylogenetic analysis, the identified novel coronavirus is named as Severe Acute 50 Respiratory Syndrome Coronavirus 2 (SARS-CoV-2), and the disease spread by SARS-CoV-2 is known as "COVID-19", declared as a pandemic by World Health Organization (WHO).(1) 51 52 Despite global massive efforts to control the outbreak of COVID-19, this pandemic is still on the rise. To date, lack of approved medicine or vaccine impede escalated the management of the 53 54 COVID-19 epidemic. In the absence of an effective treatment strategy, developing affordable screening for rapid diagnosis is critically required in the management of COVID-19.(2) According 55 to the WHO, the immediate priority is the development of point-of-care tests for the detection of 56 57 SARS-CoV-2 at an early stage with improved sensitivity.(3)

58 Currently, the COVID-19 diagnostic test falls into two categories: antibody and nucleic acid-59 based detection systems. The Developed immunoassay is rapid but inefficient for the detection 60 of the pathogen at an early stage of infection.(2) Besides, WHO does not currently recommend 61 the use of antigen-based rapid diagnostic tests for patient care, but encourages the related 62 research to improve their performance and potential diagnostic utility.(4)

Among the nucleic acid-based detection systems, the WHO considered a Real-time polymerase chain reaction (RT-PCR) based method as a gold standard for COVID-19 testing.(4) The sample-to-result time of the quantitative RT-PCR (qRT-PCR) was initially >4 hrs; however, constant efforts are underway to improve the turn-around time through automation. Besides, time-consuming process PCR based tests are expensive; they require sophisticated instruments and expertise.(2) Thus, other technologies, such as reverse transcription- loop-mediated isothermal amplification (RT-LAMP) (5) and Genome-editing (6), are being explored. These are

promising technologies; however, the expected turn-around time would still be around 1 hour
and may not be economical for mass screening, especially for the developing countries.

Nanomaterials based sensing platforms hold promise to develop rapid disease diagnostics. However, their limited use in the clinical setting is due to the need for sophisticated equipment.(7, 8) A recent attempt to use antisense-nucleotide capped gold nanoparticles for Ngene based, COVID-19 detection could be a game changer.(9) Whereas, WHO suggested that N-gene has a relatively weak analytical capability, compared to the *RdRp* gene, to detect COVID-19 infection.(10) Moreover, the authors have demonstrated COVID-19 detection in a cellular-model system, and application in human samples has yet to be shown.

79 In this study, we report for the first time gold nanoparticles (AuNPs) based rapid colorimetric assay for visual eye detection of COVID-19 RNA in human samples designed to target RdRp 80 81 specific gene target in very cost-effective manner with wide application of mass screening in field. We utilized the surface plasmon resonance property of AuNPs to detect unamplified 82 COVID-19 RNA in human samples. The ability of AuNPs to preferentially adsorb ssRNA/ssDNA 83 84 over dsDNA/dsRNA is the crucial concept of this assay. The single and double-stranded oligonucleotides have different electrostatic properties, which provide stability (via retaining 85 86 natural color) and aggregation (which causes color change) of AuNPs in solution, 87 respectively.(11)

88 Materials and Methods

Chemicals: Citrate buffer stabilized AuNPs (10 nm diameter) was purchased from Alfa Aesar
 (Thermo Fisher Scientific India Private Limited). Phosphate Buffer Saline (PBS pH 7.0) and
 NaCl were procured from Sigma Aldrich.

92 *Clinical samples:* Nasopharyngeal RNA sample from COVID- negative (n=6) and positive 93 subjects (n=7) were used. The COVID-19 testing of these subjects were done in the Indian

Council of Medical Research (ICMR, India) approved diagnostic laboratory using Tagman-94 based RT-PCR kit (Labgun, lab, Genomics.co. Ltd, Republic of Korea). The study protocol to 95 use clinical samples was approved by Institutional Human Ethics Committee SGPGIMS, 96 97 Lucknow (Ref N. PGI/BE/327/2020). The cervical DNA samples from human papilloma virus 98 (HPV) infected woman (n=2) was used as non-specific target control. The clinical Samples (cervical smear) for the HPV DNA test were processed using HPV Test Hybrid Capture® 2 99 100 protocol (QIAGEN). Samples with relative light units (RLU) /Cutoff Value ratios > 1.0 were considered as HPV positive and < 1.0 were considered as HPV negative. Only HPV positive 101 DNA samples were used during validation experiments. 102

In Vitro transcription (IVT) of SARS-CoV-2 RNA: The RNA (5ng) from confirmed COVID-19 positive human sample were first reverse transcribed with modified oligo dT primer having a T7 promoter sequence at its 5' end, and the resulting single stranded cDNA was further *In-vitro* transcribed (IVT) using HiScribe T7 Quick High Yield RNA Synthesis Kit according to the manufacturer's protocol (New England BioLabs Inc, NEB #E2050). The amplified RNA was than purified using Monarch RNA Cleanup Kit (New England BioLabs Inc.) and quantified by RNA HS reagent using Qubit system (Thermo Fisher Scientific).

Colorimetric assay for the detection of SARS-CoV-2 RNA: For colorimetric assay, reaction 110 was set up in 10 µL of reaction volume in sterile PCR tubes, containing (a) hybridization buffer 111 112 containing 80mM NaCl), (b) 0.5 µM oligo probe for RdRp genes, and (c) target RNA from positive patients or IVT RNA. The RdRp oligo probe (5'-GTGATATGGTCATGTGTGGCGG-3') 113 114 was used to specifically detect the presence of SARS-CoV-2 RNA in the assay. In parallel, to 115 measure the specificity of the assay, input genomes from different source such nasopharyngeal RNA from COVID negative subjects and HPV DNA from cervical cancer positive samples were 116 117 used as negative controls. Similarly, non-template control (NTC) was also included to measure the background reactivity. In addition, to confirm the working principle of the assay, a different 118

119 RNA template (isolated from pancreas) and pancreas specific *REG-3* (Regenerating islet-120 derived protein 3) gene oligo probe (5'- GTGCCTATGGCTCCTATTGCT-3') were used 121 separately.

The final reaction mixture with above combinations was then denatured at 95°C for 30 seconds, annealed at 60°C for 60 seconds and then cooled to room temperature for 10 minutes. Subsequently, 10 nM colloidal AuNPs (~10 nm) were added to the assay mixture and allowed to develop color for 1-2 minute.

Spectral studies and measurement of sensitivity: Absorption spectrum of the assay mixture 126 127 was recorded in the range of 300 -700 nm. The peak shift from 520 nm (known as red-shift) and 128 peak broadening after 520 nm were measured as a characteristic feature of the salt induced aggregation. Using various combinations of positive and negative controls (as discussed in the 129 130 previous section) the specificity of reaction and aggregation were compared. Assay sensitivity was determined by serially diluting the input SARS-CoV-2 RNA from both the IVT synthesized 131 and, synthetic SARS-CoV-2 control (nCov19 control kit by Applied Biosystems) ranging from 5-132 0.1ng and 1-0.1ng concentrations respectively. 133

134

135 **Result and discussion**

In this study, we report for the first time the development of a rapid and affordable RNA-based assay for the visual detection of the SARS-CoV-2 genome in human samples. In this assay, we use surface plasmon resonance property of gold nanoparticles/colloids (AuNP) and targeted the *RdRp* specific gene sequence of SARS-CoV-2. *RdRp* is essential for viral replication and has higher analytical power than E (envelope protein) and N (nucleocapsid protein) genes of SARS-CoV-2. (10) our current established assay, salt-induced aggregation and color change of the gold colloids occurs after *RdRp* oligo probe hybridizes with its specific target RNA of SARS- 143 CoV-2. With the current escalated demand of cost effective, easy and sensitive diagnostic for 144 COVID-19, the test was developed using commercially available nCoV19 synthetic DNA and 145 validated further using clinical samples from COVID-19 subjects (as confirmed using Taqman 146 based RT-PCR method, Table S1). In our study, we demonstrated a visual change in gold 147 colloid color from pink to blue when RNA samples from subjects with clinically diagnosed 148 COVID-19 infection hybridize with *RdRp* oligo probe (n=7). Simultaneously, the color remained 149 pink in SARS-CoV-2 negative samples due to the absence of hybridization (n=6).

Figure 1 illustrates the sequential schematics of process of the developed test. The color of the 150 gold colloid solution is dependent on the aggregation property of AuNPs in suspension.(12, 13) 151 152 The aggregation-induced color change can be visually monitored (by the naked eye) and quantitated through absorption spectroscopy. Generally, in aqueous solution, gold colloids 153 154 remain stabilized by the coating of negatively charged citrate ions (14) and have visible 155 appearance of pink color (Figure 1). In solution, individual particle exhibits a surface plasmon 156 resonance peak (λ max) at 520 nm (Figure 2a, Green curve). The oligonucleotide probe preferentially adsorbs on AuNPs and provides additional stability due to the addition of negative 157 158 charges.(15) The same phenomenon was observed in our experiments when RdRp oligo probe adsorbed and protected the salt-induced aggregation of colloid in the absence of target RNA 159 (NTC) (Figure 2a, Grey curve). Except for the reduced intensity of the absorption spectrum, 160 which was due to dilution of the colloid (Figure 2a, Grey curve), NTC and AuNP assays showed 161 no change? A pure colloid is pink in color (Inset of Figure 2a left vial), which turns into light pink 162 when it reacts with hybridization buffer containing oligo probe in equal volume (without target) 163 164 (Inset of Figure 2a right vial). Unlike dsDNA, inherent structural flexibility of ssDNA/RNA to 165 partially uncoil its bases, exposing them to AuNPs, generates the attractive electrostatic forces 166 causing them to allow over colloids and giving protection against electrostatic interaction 167 causing salt-induced aggregation.(15)

In this assay, SARS-CoV-2 RNA from human patients or IVT synthesized RNA was added into 168 169 hybridization buffer (containing oligo probe), followed by denaturation and annealing at 95 C (30s) and 60°C (60s), respectively. After cooling at room temperature, the gold colloid was 170 171 added into the above reaction mixture. The colloid color changes in visible range from pink to 172 blue, indicate the formation of hybridized product (Figure 2b-c, and Figure 3). Broadening of the peak with red-shift (~30 nm) was observed in the spectrum of aggregated colloids than non-173 174 aggregated, confirms the success of developed assay for detection of an unamplified target with unmodified colloids in a quick and facile way. The principle of binding oligo probe to its specific 175 target leading to change in color of the solution was independently verified using a different 176 177 template RNA (isolated from pancreas tissue) and pancreas specific gene REG3 oligo probe in a separate assay. This assay also resulted in a similar change in color and absorption spectra 178 179 as optimized earlier for SARS-CoV-2 RNA and RdRp oligo probe. It established the working 180 principle and specificity of the test (Figure S1).

181 We determined the cross-reactivity using a cervical-DNA sample from women diagnosed with 182 HPV infection (non-specific target control). No color change of gold colloids was observed with HPV DNA, indicating no hybridization, and specificity of the developed assay (Figure 3 b, right 183 184 vial). Contrary to HPV DNA-negative control and NTC, a positive control sample shows development of blue color (Figure 3 b, middle vial). Absorption spectrum (of colloids) with HPV 185 DNA-negative control exhibited characteristics similar to that of NTC, and no red-shift or peak 186 broadening as found with positive samples (Figure 3a). Cross-reactivity of the developed assay 187 188 with other respiratory viruses is warranted. However, we do not anticipate the same as the test 189 utilizes the detection of the RdRp gene of the SARS-CoV-2 virus. The oligo probe sequence used in our assay is not complementary to any human mRNAs and other members of the SARS 190 family, as verified by BLAST using the NCBI database. (10) 191

192 IVT synthesized SARS-CoV-2 RNA was used to test the sensitivity of the developed assay.
193 RNA ranging from 0.1 to 5ng resulted in a gradual change in colloid color from light pink to blue
194 (Figure 4b). The color difference at 0.1 ng, compared to NTC, is barely visible with naked eyes
195 (Figure 4 b extreme left vial). However, the absorption spectra of colloids show a clear red-shift
196 with peak broadening up till 0.5 ng target RNA (Figure 4a).

We determined the assay sensitivity using different dilutions (1-0.1ng) of PCR-amplified 197 198 synthetic DNA (positive control, with nCoV19 control kit). Similar to IVT synthesized RNA, a decreasing amount of DNA (from 1 to 0.1 ng) show a gradual change in colloid color from blue 199 200 to light pink (left to right, Fig. 4d). A clear red-shift with peak broadening reflects in the absorption spectra of colloids recorded with control DNA, compared to NTC (Fig. 4c). 201 Accordingly, positive control showed a clear visual demarcation up to 0.5 ng amount, compared 202 203 to NTC (Fig. 4d). A similar recent approach utilizes thiol capped gold nanoparticles to detect N-204 gene of the SARS-CoV-2 gene in a cellular system.(9) However, for the detection of SARS-CoV-2 RNA in human samples, the N-gene has reportedly inferior analytical power than the 205 RdRp gene.(10) Thus, it would be essential to know the assay's performance, developed by 206 207 Moitra et al., with clinical samples.

208 Conclusions

We have successfully developed an affordable gold nanoparticles-based colorimetric test for the rapid detection of SARS-CoV-2 RNA in humans. The assay can detect up to 0.5 ng of SARS-CoV-2 RNA. The turnaround time of our assay is less than 30 minutes. Moreover, the developed test will be helpful for mass screening, as it does not require sophisticated equipment. Validation of the developed assay using a large number of clinical samples is underway to determine the diagnostic accuracy, including sensitivity, specificity, false-positive and negative rate.

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- 221 Contributions: ST, VK; conceived the idea; ST, VK, SM, RS; performed the experiments, JY,
- 222 UG; provided clinical samples and the diagnosis; TK, LS and SKV; provided critical comments on
- the manuscript draft; ST, VK and SM; drafted the manuscript.
- 224

225 Competing Interest

226 The authors declare no conflict of interest.

227 Supplementary Files

- 228 Supplementary information supporting the finding of this study is available in this article as 229 supporting information.
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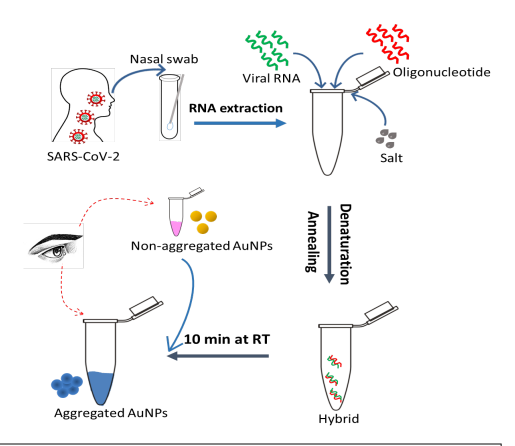
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284 Figure 1.

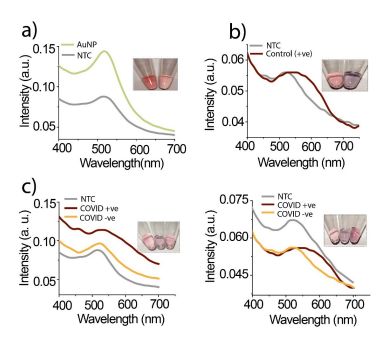


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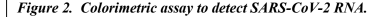
Figure 1. Schematic representation of the assay for the visual detection of SARS-CoV-2 RNA

Schematic illustrates the assay flow to detect *RdRp* (RNA dependent RNA polymerase) gene sequence of SARS-CoV-2 in the nasopharyngeal RNA sample from subject clinically diagnosed with nCOVID infection (**positive control**). Hybridization buffer with *RdRp* oligo probe (forward) was mixed with RNA sample. The reaction mixture was denatured at 95°C for 30 seconds, followed by annealing at 60° C for 60 seconds. After annealing, the tube was kept at room temperature for 10 minutes before colloidal AuNPs were added. A pure colloid is pink in color. It turns blue in the vial containing RNA sample from **positive control** due to salt-induced aggregation upon successful hybridization between oligo probe and target RNA. The color remained pink in the absence of target RNA, or the presence of a non-specific target.

286 Figure 2.



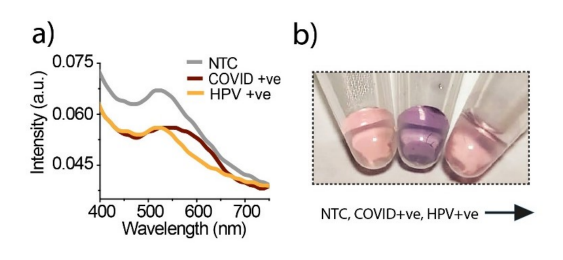
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(a) Comparative absorption spectra of unmodified AuNPs (Green curve) and oligo probe stabilized AuNPs i.e., NTC (Grey curve). Both spectra are exhibiting characteristics absorption peak at λmax 520 nm, however, the reduced peak intensity in NTC is due to dilution of colloid solution. In NTC, no red-shift in peak position, approve the stabilizing property of single-stranded oligo probe, Optical images of gold colloids (left vial) and NTC (right vial) are shown in the inset. **(b)** Comparative absorption spectra of NTC (Grey curve) and positive control i.e., nasopharyngeal RNA sample from subject clinically diagnosed with nCOVID infection (Brown curve). Broadening of the peak as well as red-shift in peak position confirm the salt-induced aggregation of AuNP due to successful hybrid formation in control. Optical images shown in inset demonstrate the evident change in the color of the solution from pink to blue in the control vial (right) while no change in color of NTC vial (left). **(c)** Representative absorption spectra, and in the inset shows optical images comparing assay performed with NTC (Grey curves, left vial), RNA from clinically diagnosed nCOVID infected subjects (Brown curve, middle vial), and RNA from subjects without nCOVID infection (Yellow curve, right vial). Samples from a total of seven infected and six uninfected individuals were analyzed.

288 **Figure 3**.

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Figure 3. Specificity of developed assay to detect SARS-CoV-2 RNA.

Nasopharyngeal-RNA from nCOVID infected subject (positive control), and a cervical DNA sample from Human papillomavirus (HPV, non-specific target control) infected women were tested. **(a)** Comparative absorption spectra for no target control (NTC, Grey curve), positive control (Brown curve), and nonspecific target control (HPV, Yellow curve). In the positive control, the broadening of the peak and redshift in peak position (Brown curve) confirmed the salt-induced aggregation upon successful hybrid formation. **(b)** Optical pictures demonstrate that the colloid color from pink to blue changes only in the vial with the positive control (middle vial), while the color remained pink in the vial with NTC (left vial) or HPV (right vial).

293 Figure 4.

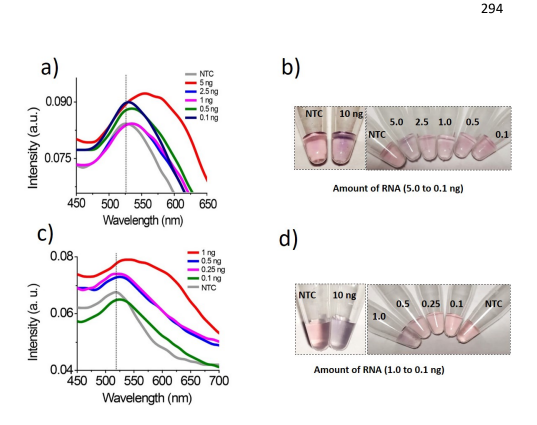


Figure 4. Sensitivity of developed assay to detect SARS-CoV-2 RNA.

Assay sensitivity was determined using different concentrations (concentration are in ng) of IVT synthesized SARS-CoV-2 RNA (a-b), or nCOVID synthetic DNA (c-d). Absorption spectra (a and c) corresponding to aggregated colloids exhibit a clear red-shift in peak with broadening, indicating successful hybridization. Optical pictures (b &d) of the assay performed to demonstrate the color change, an extra pair of vials (on the left) to show color change with a higher amount (10ng) of target nucleic acid for the reference

297 Supporting information

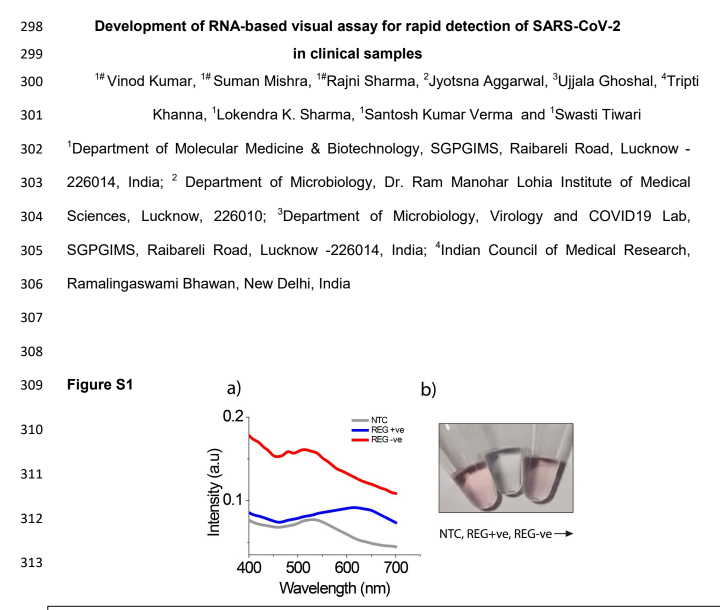


Figure S1. The assay was performed using an oligo probe specific for regenerating islet-derived (REG 3). The assay was containing RNA extracted from pancreatic tissue (positive control), placental RNA (negative control), or no RNA (NTC). (a) Comparative absorption spectra of NTC (Grey curve), positive control (Blue curve), and negative control (red curve). In the positive control, the broadening of the peak and red-shift in peak position (Blue curve) confirm the salt-induced aggregation upon successful hybrid formation. (b) Optical pictures demonstrate that the colloid color from pink to blue changes only in the vial with the positive control (middle vial), while the color remained pink in the vial with NTC (left vial) or negative control (non-specific target, right vial).

314 Table S1

315

316	Patient ID	Age in Years (Y)/ Gender	nCOVID19 infection	E gene Ct value	RdRp Ct Value
317	S1	18 Y / male	Positive	25.45	25.30
318	S2	18Y / male	Positive	28.37	29.20
510	S3	11 Y /female	Negative	-	-
319	R1	1 Y/Female	Positive	22.66	21.20
220	R2	28 Y/Female	Negative	-	-
320	R3	35 Y/Male	Negative	-	-
321	R4	66 Y/Female	Positive	22.81	24.59
222	R5	18 Y/Male	Positive	23.16	22.66
322	R6	14 Y/Male	Negative	-	-
323	R7	70 Y/Female	Negative	-	-
324	R8	44 Y/Male	Positive	29.64	25.56
524	R9	35Y /Male	Negative	-	-
325	R10	9 Y/Female	Positive	28.69	29.70

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Table S1. The table shows the clinical diagnosis of the subjects based on Taqman RT-PCR analysis. RNA extracted from the nasopharyngeal samples were analyzed in the study.

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