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Prospective evaluation of ID NOW COVID-19 assay used as point-of-care test in an emergency department

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ABSTRACT

Background: Rapid testing for COVID-19 has been clearly identified as an essential component of the strategy to control the SARS-CoV-2 epidemic, worldwide. The ID NOW COVID-19 assay is a simple, user-friendly, rapid molecular biology test based on nicking and extension amplification reaction (NEAR).

Objectives: The aim of this study was to evaluate the ID NOW COVID-19 assay when used as a point-of-care test (POCT) in our Emergency Department (ED).

Type of study: This prospective study enrolled 395 consecutive patients; paired nasopharyngeal swabs were collected from each study participant. The first swab was tested with the ID NOW COVID-19 assay at the point-of-care by ED nurses. The second swab was diluted in viral transport medium (VTM) and sent to the clinical microbiology department for analysis by both the RT-PCR Simplexa test COVID-19 Direct assay as the study reference method, and the ID NOW COVID-19 assay performed in the laboratory.

Results: Nasopharyngeal swabs directly tested with the ID NOW COVID-19 assay yielded a sensitivity, specificity, PPV and NPV of 98.0%, 97.5%, 96.2% and 98.7%, respectively, in comparison with the RT-PCR study reference assay. When the ID NOW COVID-19 assay was performed in the laboratory using the VTM samples, the sensitivity decreased to 62.5% and the NPV to 79.7%. Three false negative test results were reported with the ID NOW COVID-19 assay when performed using undiluted swabs directly in the ED; these results were obtained from patients with elevated CT values (> 30).

Conclusion: We demonstrated that the ID NOW COVID-19 assay, performed as a point of care test in the ED using dry swabs, provides a rapid and reliable alternative to laboratory-based RT-PCR methods.

1. Introduction

Coronavirus disease 2019 (COVID-19), caused by SARS-CoV-2, first appeared in China and then spread worldwide [1]. The primary goal of the epidemic containment of COVID-19 is to reduce the infection transmission in the population by reducing the number of susceptible persons or by reducing the basic reproductive number (R0).

To date, the reference testing method is the real-time reverse transcriptase-PCR (RT-PCR) similar to that developed for the diagnosis of SARS-CoV in early 2000s [2]. However, due to the rapid spreading of the SARS-CoV-2 pandemic and the limited capacities of molecular testing at the laboratory level, the concept of molecular testing in point-of-care setting such as in an Emergency Department (ED) appears to be useful to manage suspected infection cases. Indeed, the urgent need for increased testing for COVID-19 has been clearly identified as an essential component of the strategy to control the epidemic, worldwide.

The ID NOW COVID-19 assay is a simple, user-friendly, rapid molecular biology test and do not required specific equipment molecular biology, allowing point-of-care testing. This molecular isothermal assay is based on nicking enzyme amplification reaction (NEAR) technology that targets the RdRP gene [3,4] with a manufacturer’s claimed limit of detection (LOD) of 125 genome equivalents/mL. Results are provided in

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approximately 15 min, including upfront 3 min of sample elution buffer
warm-up time. The assay run time on the ID NOW platform was within 2
to 5 min for a positive call in the early callout mode as opposed to less
than 13 min on the full-callout mode for a negative result. To date, the
reported analytical performance of the ID NOW COVID-19 test in the
literature reviewed has been variable [5–7]. This could be related to the
conditions under which the samples were collected. The manufacturer’s
instructions state that this test is valid only on direct swab and not after
discharged in viral transport medium (VTM) [3]. The objective of this
work was first to evaluate the analytical performances of the ID NOW
COVID-19 assay performed by ED nurses in point-of-care in comparison
with the reference test RT-PCR Simplexa COVID-19 Direct assay (Dia-
Sorin) currently in use in our laboratory. The second objective was to
evaluate the effect of the transport medium VTM on the performance of
the test.

2. Materials and methods

2.1. Type of study

This prospective study was conducted between October 22 and
December 22, 2020 in the ED of Groupe Hospitalier Paris Saint-Joseph,
France. Consecutive adults presenting with clinical suspicion of COVID-
19 assessed by the attending emergency physician and submitted to
diagnostic tests for SARS-CoV-2 were eligible. The main clinical and
laboratory data were collected in an electronic Case Report Form (e-
CRF).

All adult patients consenting to the participation in the study and to
the use of biological samples were included. Patients were excluded
when samples missed or inadequate.

2.2. Clinical specimens

Paired nasopharyngeal swabs were collected with a flexible naso-
pharyngeal flocked swabs from patients having a clinical suspicion of
COVID-19 by the attending nurse in the ED. Swabs were sent to the
microbiology laboratory without prior assignment to either technique.
The lack of identification of the order of collection and the size of the
study ensures that the swabs were randomly distributed.

- One swab was directly used for the ID NOW COVID-19 assay in point-
of-care test by ED trained nurses, previously trained and certified for
use it. Training on the use of ID NOW was provided by qualified
experts. Proficiency tests were conducted with ED nurses prior to study
initiation. Testing was conducted according to instructions for
use for ID NOW.
- The second swab collected, was then discharged in 3-ml viral
transport medium (VTM) (Labo Moderne, Gennevilliers, France) and
sent to the central microbiology laboratory for analysis.

Specimens were stored at 4 °C up to a day, if all testing could not be
completed on the same day. In case of discrepancies, samples were ali-
quoted and frozen at −80 °C for confirmatory testing.

2.3. Testing methods

The ID NOW COVID-19 assay (Abbott, Chicago, Il, USA) is an
isothermal nucleic acid amplification-based. The assay was performed
directly from the dry swab in the ED or by transferring 200 microliters of
VTM to elution buffer in the sample base and then mixed for 10 seconds
per instructions for use at the microbiology laboratory.

The Simplexa COVID-19 direct assay (DiaSorin, Saluggia, Italy) was
chosen as RT-PCR reference test and was performed with the DiaSorin
LIAISON® MDX according to the manufacturer’s instructions for use. A
50 μl volume of Simplexa COVID-19 Direct kit reaction mix (MOL4150)
was added to the “R” well of the 8-well direct amplification disc
followed by addition of 50 μl of non extracted nasopharyngeal swab
sample to the “SAMPLE” well. Fluorescent probes are used together with
corresponding forward and reverse primers to amplify two different
regions of the SARS-CoV-2 genome: ORFlab and S gene. Data collection
and analysis were performed with LIAISON® MDX Studio software. CT
values were collected from MDX software.

2.4. Confirmation of discrepant results

In case of discrepancy, a control was performed using an aliquot from
VTM previously stored at −80 °C for quantitative RT-PCR following the
protocol established by the National Reference Center for Respiratory
Viruses (NRC) at Institut Pasteur, Paris, France [8,9]

2.5. Statistical analysis

The percentages were calculated based on documented data (missing
data were excluded from the percentages). Inter-group comparisons
were made using the Mann-Whitney test for quantitative variables and
the Fisher exact test for qualitative variables. Statistical analysis was
done with StatView software (version 5.0). All tests were two-tailed and
p values less than 0.05 (calculated by χ² test, Student’s t test, or Mann-
Whitney test) were considered significant.

2.6. Ethical statement

This study followed the Standards for Reporting Diagnostic Accuracy
study (STARD) guidelines and was previously approved by the local
clinical ethic committee board N° IRB 2020-A02758–31. Informed oral
consent for participation was obtained from each participant, in accor-
dance with French law.

3. Results

A total of 406 patients were enrolled in the study, mean age was 71
years and M/F sex ratio = 1.12. Among them, 11 were secondary
excluded due to a lack of adequate specimen samples (Fig. 1: Study Flow
Chart). A total of 395 patients were eligible for study inclusion. Among
these, 154 patients (39%) were diagnosed with SARS-CoV-2 infection
(Fig. 1).

3.1. Performances of the ID NOW COVID-19 assay compared to
reference RT-PCR assay

POCT in the ED with nasopharyngeal swabs directly tested using the
ID NOW COVID-19 assay showed sensitivity, specificity, positive pre-
dictive value (PPV), and negative predictive value (NPV) of 98.0%,
97.5%, 96.2% and 98.7%, respectively. When the ID NOW COVID-19
assay was performed in the laboratory from the swab previously dis-
charged in VTM, the sensitivity dropped to 62.5% and the NPV to 79.7%
(Table 1). Three false negative test results were reported with the ID
NOW COVID-19 assay when performed using undiluted swabs directly
in the ED; these results were obtained from patients with elevated CT
values (> 30). Six false positive results were observed with ID NOW
COVID-19 assay performed in emergency department and no false
positive results were reported in microbiology laboratory testing.

3.2. Comparison of Ct values among positive samples detected by direct ID
NOW COVID-19 assay and samples performed on VTM by ID NOW
COVID-19 assay

The median cycle threshold (CT) for positive sample by direct ID
NOW COVID-19 assay and ID NOW COVID-19 assay performed on VTM
were 17.9 (interquartile range [IQR] [15.5–21.3]) whereas for positive
sample only by direct ID NOW COVID-19 assay median (CT) were 30.7
[28.2–32.4]; p < 0.001 (Fig. 2). Overall agreement between direct ID
We observed nine discrepancies between the results with the ID NOW COVID-19 assay performed directly in the ED and the RT-PCR method by Simplexa at the central laboratory (Table 1). An aliquot from VTM were sent to the National Reference Center for respiratory viruses at Institut Pasteur, Paris. Finally, we reported three samples (ID NOW COVID-19 assay negative, positive by RT-PCR) and six samples (ID NOW COVID-19 assay positive, control negative by RT-PCR).

With the three false negative samples, we made two observations: either a lack of detection of a low viral load SARS-CoV-2 infection (n = 2) or a previous history of SARS-CoV-2 infection (n = 1). Two patients presented with symptoms suggestive of SARS-CoV-2 infection. One had pneumonia associated with a biological inflammatory syndrome and lymphopenia. The other patient had acute viral gastroenteritis. The Simplexa MDX RT-PCR assay was performed within 24 h of collection and positive result with a high CT in both cases (> 30), indicating a low viral load. The third patient had a history of COVID-19 six weeks earlier in a pauci-symptomatic form. She was being managed in the ED for a bowel obstruction and had no symptoms of COVID-19. Simplexa MDX RT-PCR control and National Reference Center for Respiratory Viruses (NRC) control confirmed the persistence of low viral load with high CT (CT > 30) related to the previous infection and were not detected by the Abbott ID NOW assay as it is less sensitive. Heterogeneity of viral load between the two swabs may also explain these discrepancies.

Among the six false-positive swabs, we were also able to make two observations: known previous SARS-CoV-2 infections (n = 3) and early forms of SARS-CoV-2 infections (n = 3). Among the three patients with a history of COVID-19, two had been hospitalized for reasons unrelated to and distant from COVID-19 (6 weeks and 8 months after). The third had been hospitalized three weeks after the diagnosis of COVID-19 for an episode of dyspnea. Thoracic angioscanner found minimal (10–25%) sequential lung damage with no pulmonary embolism. After investigation, we have no confirmation by a follow-up of the detection of SARS-CoV-2 for the 3 patients concerned but we have the clinico-scannographic confirmation during the hospitalization with oxygen requirement, typical symptoms and evocative CT-scan. Simplexa, and

Fig. 1. Study flowchart. The arrow indicate the study flow, up to the final classification of 154 patients with the infection and without the infection.

### Table 1
Analytical performances of the COVID-19 ID NOW test assay performed in emergency department and ID Now test in microbiology laboratory compared to reference RT-PCR assay.

|                     | Detected | Not detected | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) |
|---------------------|----------|--------------|----------------|----------------|---------|---------|
| ID Now test performed in ER |          |              |                |                |         |         |
| Detected            | 151      | 6            | 98.0           | 97.5           | 96.2    | 98.7    |
| Not detected        | 3        | 235          |                |                |         |         |
| ID Now test in microbiology laboratory |          |              |                |                |         |         |
| Detected            | 100      | 0            | 62.5           | 100            | 100     |         |
| Not detected        | 60       | 235          |                |                | 99.7    |         |

Fig. 2. Comparison of CT values among positive samples detected by direct ID NOW COVID-19 assay and ID NOW COVID-19 assay performed on VTM. CT value differences were compared by using the Kruskal-Wallis test. VTM: viral transport medium.
NRC Pasteur RT-PCR controls showed negative results which may be explained by heterogeneity of loading between the first and second nasal swab. Among the three patients with no history of COVID-19, timing context and associated symptoms were consistent with frequent modes of COVID-19 disclosure in the elderly: fall, anorexia, weakness and delirium. The discrepancy could probably result from the lower quality of the second swab, which remains an invasive sample. We hypothesize that these cases were SARS-CoV-2 infections not detected by the RT-PCR controls because of a viral load defect on the second swab. Nevertheless, given the lack of evidence to specifically identify that these cases were early SARS-CoV-2 infections, other hypotheses could explain the false positives, notably cross-contamination and assay specificity issues.

4. Discussion

Rapid and accurate detection of SARS-CoV-2 is essential to ensure early and appropriate patient management, outbreak containment, and to better follow the global epidemiology of the virus. Laboratory testing is mainly based on amplification and detection of viral gene sequences in upper respiratory tract specimens. The results of these analyses are often lengthy (1–4 h) and difficult to manage with a large flow of patients in the ED and lead to a risk of overcrowding in these units. As reported by Alter et al. [10] and our team [11], point-of-care test (POCT) instruments in ED produce lab results with rapid turnaround times.

This study demonstrated that the detection of SARS-CoV-2 using rapid isothermal nucleic acid amplification assay performed by nurses directly at the emergency room, 24 h-a-day, 7 days-a-week was both feasible and reliable. The sensitivity and specificity of the ID NOW system for the detection of SARS-CoV-2 was higher than 98% even when performed by a trained professional (in this case a nurse) without direct laboratory supervision. Less than 3.8% of tests failed were observed during this study. As reported by Kanwar et al. [12], the invalid rate is an important consideration when selecting an assay for clinical use. Our experience with the ID NOW platform for the diagnosis of seasonal influenza showed satisfactory results [13] which allowed implementation POCT in our ED [11]. This experience has enabled the rapid implementation of a POCT’s strategy in ED for the diagnosis of COVID-19.

We have to keep in mind that identification of the virus is only a part of clinical diagnosis and management. Although our results are important, further studies are necessary to assess the clinical impact of ID NOW COVID-19 assay at triage in order to evaluate its effect on decision-making and the prescription of complementary examinations. POCT is only useful if it has an impact on treatment or healthcare organization. Some studies conducted during influenza outbreaks have shown a positive impact on medical costs [14,15], rapid antiviral treatment [16] and reduced length of stay [11], especially for patients with a negative result, highlighting the importance of tests that have a high negative predictive value. However, we demonstrated the significant decrease in sensitivity of the ID NOW COVID-19 assay when performed using diluted swabs in VTM. Thus, we strongly recommend to perform immediately the test on fresh swab only. Dilution of the nasopharyngeal swab specimen in VTM below the lower limit of detection for the assay increases the risk of false negative results. This probably explains the better sensitivity of the test that we find compared to the results described by Mitchell and George [17].

Similarly, it is interesting to note that in the study by Basu et al. [18] 101 paired nasal samples were tested on direct swabs and VTM transport medium. Fifteen samples were positive with the Cepheid GeneXpert test and only 10 samples were positive by the Abbott ID NOW test [19]. Thus 5/15 were false negatives. However, it is important to emphasize that in this study the 5 negative samples had high Ct values >33 and even very high Ct values for 3 samples (>40 on one of two nucleic acid targets). The clinical significance of these very low viral load levels is largely unknown. Nevertheless, as reported by Basu et al. ID NOW COVID-19 assay has some utility as a rapid rule-in test for samples at high viral load [18].

Furthermore, Abbott claim a sensitivity of 125 copies /ml for the ID NOW assay [3,4]. However the evaluation by Department of Health Social & Care (DHSC) showed the assay starts to drop out at PCR Ct >30 values in comparison with reference quantitative PCR assays [20]. The publication by Zhen et al. [7] reported a LOD for the Abbott ID NOW assay of 20,000 copies /ml. Discordant samples in this study negative by the Abbott ID NOW assay and positive by the reference RT-PCR method had RT-PCR Ct values between 30 and 38. Based on this, it seems unlikely the claimed LOD by Abbott of 125 copies /ml is correct. Data published by Abbott also indicate poorer sensitivity at higher comparative PCR Ct values >33.

In the case of invalid or negative result with clinical signs suggestive of SARS-CoV-2 infection, the physician should perform an RT-PCR confirmatory test. However, once the swab is discharged into the sample receiver, if the result is invalid, it is no longer usable for central laboratory test.

As described by Mitchell and George [17] the potential for error and cross contamination from the multiple cartridge transfers and manipulations and possible biosafety concerns while swabbing the swab to resuspend material are contingencies that the operator should be cautioned about during training. Nevertheless, no cross-contamination was observed in Mitchell’s study, nor any false positives (100% specificity). Therefore, for the 6 false positive samples, the hypothesis of a viral load defect on the second swab is the most plausible. This is supported by the fact that no false positives were observed on the viral transport medium VTM by both the IDNOW microbiology test and the Simplexa RT-PCR.

Our study has several limitations. First, the ID NOW COVID-19 assay itself has only one target, the RdRP gene. To date, the published isothermal amplification technology tests did not evaluate SARS-CoV-2 using two target genes [21]. Second, this work was conducted during an epidemic peak and so the results should be viewed in the context of the conditions during such a period. Positive predictive values decrease when tests are performed with a lower prevalence of the disease, thus the results cannot be extrapolated to a lower prevalence period. Third, this study was conducted on the relatively small number of samples in a single care center setting. Therefore, the results may be specific to the population of this center and may not be applicable to another patient group such as children for example. Use of rapid diagnostic test at triage in other care settings would depend on the organization of each ED and, importantly, on the availability of compliant and trained staff.

5. Conclusion

With large numbers of patients being admitted to hospital, putting tremendous pressure on health care systems, there is an urgent demand for a user friendly, rapid, simple and sensitive POCT assay. Such tests could be used at hospitals to facilitate faster detection of SARS-CoV-2, which can reduce or avoid further spread and ensure appropriate patient management. This is the first prospective evaluation of ID NOW COVID-19 assay in “real life” in ED and we showed ID NOW COVID-19 assay used as POCT in an ED setting provides a rapid and reliable alternative to laboratory-based RT-PCR methods.

Declaration of Competing Interest

We have no conflicts of interest to declare.

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