Evaluation of sensitivity and specificity as COVID–19 screening method

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ABSTRACT

Coronavirus disease 2019 (COVID-19) is caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). Antibody rapid test is one of the COVID-19 screening tests that can be used in the community. The accuracy of the rapid antibody methods needs to be appropriately assessed, it is necessary to carry out a diagnostic accuracy study using a pairwise sensitivity and specificity analysis. This research aimed to assess the sensitivity and specificity of COVID-19 rapid tests, also assesses positive predictive value (PPV) and negative predictive value (NPV) of the rapid antibody test as a method of screening for COVID-19 in Sleman Regency, Indonesia. In total, 118 respondents who have contact with COVID-19 patients and have symptoms were enrolled in this study. The study was conducted on 118 patients who met the close contact criteria were conducted a rapid antibody test. 64.41% patients were reactive. Real-time polymerase chain reaction (RT PCR) as a gold standard was also carried out for all patients and 63.56% affirmed positive for COVID-19. The sensitivity value was 97.33%, and the specificity value was 93.02%, while the positive predictive value (NPP) was 96.05%, and the negative predictive value (NPN) was 95.24%. These results meet the minimum recommendations for the screening method.

Keywords: Antibody rapid test, COVID-19, Screening, Sensitivity, Specificity

1. INTRODUCTION

Coronavirus disease (COVID-19) is caused by the severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). SARS-CoV-2 transmission occurs from symptomatic patients through droplets that come out when coughing or sneezing. Meanwhile, cases related to transmission from asymptomatic patients generally have a history of close contact with COVID-19 patients [1]. The first cases of COVID-19 were reported in Indonesia on March 2, 2020, totaling two cases. Data shows that there are 1,528 confirmed cases and 136 deaths in March 2020 [2]. Meanwhile, the COVID-19 case in Sleman Regency, Indonesia was first recorded on March 18, 2020. Based on data from the Sleman Regency Health Service, on July 31, 2020, there were 233 cases, with eight of them dying, so the mortality rate reached 3.43% [3].

The fact that the entire world has been exposed to COVID-19 is causing the economy to disrupt. Therefore to re-open economic activity, health care encourages antibody tests for screening COVID-19 in the community [4]. The rapid test is use as a choice because it can detect immunoglobulin M (IgM) as a form of the body’s defense response against viral infections. It also detects Immunoglobulin G (IgG), which is an immunological and immune memory. The process of forming IgG and IgM antibodies as a result of infection
with SARS COV-2 that causes COVID-19 is an essential indication in this test [5]. In America, people are scrambling to carry out rapid antibody tests to escape lockdown [6]. But this strategy means nothing if the test result can’t be trusted [6] therefore, the accuracy of antibody test is essential. Currently, the US Food and Drug Administration have given commercial test manufacturers authorization for COVID-19 antibody test. Sensitivity and specificity are required before their use in clinical practice [4].

The process of forming IgG and IgM antibodies as a result of infection with SARS COV-2 that causes COVID-19 is an important indication in this test [5]. In general, IgM is produced earlier, followed by IgG production. However, studies of SARS COV-2 have shown that IgM and IgG development often occur at the same time [7]. Most of the cases obtained do not show clinical manifestations or asymptomatic. Therefore, the rapid antibody test is expected to have a high sensitivity and specificity level so that the examination results are accurate. The accuracy of the rapid antibody methods needs to be appropriately assessed. Sensitivity for instance depends on the method itself [8] and the timing exposure and onset of symptoms [9]. To evaluate the rapid test method, it is necessary to carry out a diagnostic accuracy study using a pairwise sensitivity and specificity analysis [10]. Sensitivity can describe the probability of measuring the likelihood for a rapid test to pick up the presence of disease; alternatively, a true positive is recorded when a procedure reflects the presence of the pathogen in a contaminated sample. Meanwhile, Furthermore, we define specificity as the probability of measuring the likelihood for a test to pick up the absence of a disease/pathogen, alternatively, a true negative is recorded when a procedure reflects the absence of a pathogen when the sample is not contaminated [11]-[13]. False-negative result may cause COVID-19 transmission in community, false-positif may impact with patient psychological condition, and patient may suffer from quarantined at home.

The CDC recommends three approaches for choosing and optimizing antibody tests. First, a population with more than 5% prevalence of COVID-19 and should choose an antibody test with high specificity [4]. Second antibody test use for any person who previously was exposed to COVID-19 and encourage patient with positive for COVID-19 to be tested with antibody test [16]. Sleman District Health-Office effort to screen for COVID-19 in the community uses the IgM and IgG Rapid Test or antibody rapid test. One study revealed that the rapid test is the right choice because it is easier to use for COVID-19 screening in the community. The prevalence of COVID-19 cases in Sleman is 8.9%, and one-off screening policy in Sleman is close contact cases and suspected are given antibody tests. The Centers for Disease Control (CDC) recommend that rapid antibodies be used in areas with a prevalence of more than 5% and given to people who have had contact with or have been exposed to COVID-19. Therefore, the researchers intend to assess the sensitivity and specificity of the antibody Rapid Test as a method of screening for COVID-19 in the Sleman Regency.

2. RESEARCH METHOD

The respondents of this study were 118 people of Sleman Regency, Indonesia who received a rapid test antibody and met the inclusion and exclusion criteria. The inclusion criteria are: i) residents in the Sleman Regency; ii) receive a rapid antibody test by the Health Office of Sleman Regency; iii) conducted on people who meet the criteria for close contact/suspect/COVID-19; iv) the rapid test kit used is in the list of recommendations government. The primary objective of this study was to assess the sensitivity, specificity, and predictive value of the antibody test. Data were analyzed using univariate analysis, then further analysis was carried out on the test results for IgM and IgG based on the results of the sensitivity and specificity calculations. Furthermore, an analysis is followed based on the results of the estimate of positive predictive value and negative predictive value. The protocol of this study was approved by the ethics commission of Ahmad Dahlan University, Yogyakarta, Indonesia number SKEP/041/KEP/VI/2020

3. RESULTS AND DISCUSSION
3.1. Result

The study was carried out in August 2020. The study was conducted on 118 patients who had been in contact with confirmed COVID 19. 91.5% of the 118 patients experienced various symptoms, a rapid test was carried out on all patient and the results showed that 64.41% were reactive. Real-time polymerase chain reaction (RT PCR) was carried out for all patients for diagnosis with 63.56% confirmed positive for COVID-19. The 118 samples were analyzed consist of 100% female samples 59.32% of the respondents' age is > 15 years old and 90.68% respondent's work status is not working. The result of antibody test and medical record patient were reviewed to conclude. Characteristics of the respondents can be seen in Table 1. The results of antibody tests conducted on respondents were as many as 76 (64.41%) people showing reactive results. Meanwhile in Table 2 shows the result of the RT PCR examination the results were 75 (63.56%) positive and 43 negatives (35.59%).

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Table 1. Characteristics of respondents, symptoms, and antibody rapid test results

| Characteristics | Symptoms of COVID-19 experienced |
|-----------------|---------------------------------|
| Age             | Asymptomatic | 1 symptom | 2 symptoms | 3 symptoms | > 3 symptoms | Total |
| ≤15 years       | 0 | 17 | 24 | 7 | 1 | 49 (41.53%) |
| > 15 years      | 10 | 9 | 24 | 13 | 13 | 69 (58.47%) |

Gender

|        | Man | Woman |
|--------|-----|-------|
| 0      | 0   | 10    |

Employment

|          | Unemployed | Employed |
|----------|------------|----------|
| 0        | 25         | 0        |

Rapid test results

|        | Reactive | Non-reactive |
|--------|----------|--------------|
| 2      | 12       | 33           |

Table 2. The results of the rapid antibody test and RT PCR test

| Check up result | Total | Percentage |
|-----------------|-------|------------|
| Antibody rapid test |       |            |
| Reactive         | 76    | 64.41%     |
| Non-reactive     | 42    | 35.59%     |
| Total            | 118   | 100%       |
| RT PCR test      |       |            |
| Negative         | 43    | 36.44%     |
| Positive         | 75    | 63.56%     |
| Total            | 118   | 100%       |

Furthermore, an assessment of all medical record samples with symptoms was carried out. Table 3 shows that the symptoms experienced by most of the respondents were not being able to smell as many as 75 people (63.56%). While the least symptoms experienced by respondents included bitter tongue, difficulty defecating, headaches, and earaches, each experienced by one person (0.85%). Based on the data that has been collected, the calculation of sensitivity, specificity, and predictive value is then performed. The results of data processing are summarized in Table 4. The results are sensitivity value was 97.33% and the specificity value was 93.02%, while the positive predictive value (NPP) was 96.05% and the negative predictive value (NPN) was 95.24%.

Table 3. Frequency distribution of antibody rapid test results based on symptoms experienced

| Symptoms experienced | Frequency |
|----------------------|-----------|
| Can not smell        | Yes 75 (63.56%) | No 43 (36.44%) |
| Could not tell the taste | Yes 55 (46.61%) | No 63 (53.39%) |
| Cold                 | Yes 26 (22.03%) | No 92 (77.97%) |
| Cough                | Yes 32 (27.12%) | No 86 (72.88%) |
| Red eye              | Yes 3 (2.54%)   | No 115 (97.46%)|
| Bitter tongue        | Yes 1 (0.85%)   | No 117 (99.15%)|
| Fever                | Yes 14 (11.86%) | No 104 (88.14%)|
| Migraine             | Yes 3 (2.54%)   | No 115 (97.46%)|
| Dizzy                | Yes 17 (14.41%) | No 101 (85.59%)|
| Nose sore/sore       | Yes 2 (1.69%)   | No 116 (98.31%)|
| Sore throat          | Yes 3 (2.54%)   | No 115 (97.46%)|
| Difficult to defecate| Yes 1 (0.85%)   | No 117 (99.15%)|
| Headache             | Yes 1 (0.85%)   | No 117 (99.15%)|
| Inflammation         | Yes 3 (2.54%)   | No 115 (97.46%)|
| Ear pain             | Yes 1 (0.85%)   | No 117 (99.15%)|
| Nosebleed            | Yes 2 (1.69%)   | No 116 (98.31%)|
| Chest tightness/shortness of breath | Yes 3 (2.54%) | No 115 (97.46%) |

Table 4. Sensitivity and specificity of antibody rapid test

| Results of the screening (rapid antibody test) | RT PCR results | Total | Sensitivity (%) | Specificity (%) | Positive predicted value | Negative predicted value |
|-----------------------------------------------|----------------|-------|-----------------|-----------------|--------------------------|--------------------------|
| Reactive                                      | Positive       | 73    | 3               | 97.33           | 93.02                    | 96.05                    | 95.24                    |
| Non-reactive                                  | Negative       | 40    | 42              |                 |                          |                          |                          |
| Total                                         |                | 75    | 43              |                 |                          |                          |                          |
Ability of the rapid test assessment is carried out using the area under the curve (AUC). The wider the AUC, the better the ability of a test to detect a disease. The ability of a test is declared well if AUC ≥ 0.7. The result is 0.1968 this means that the ability of the rapid test antibody is good for detecting COVID-19. The AUC rapid test is shown in Figure 1.

![Figure 1. The AUC](image)

### 3.2. Discussion

In this study we evaluated rapid test for screening of COVID–19 in Sleman Regency. Samples from COVID-19 cases obtained during April 2020 who’s qualify as close contact and confirmed by PCR were used as gold standard. The 55 people (75.33%) with symptom are >15 years old, these results are the same as the findings of a study in Beijing where the majority of young adults were 77% [1].

Clinical manifestations of COVID-19 cases include no symptoms (asymptomatic), mild symptoms, pneumonia, severe pneumonia, acute respiratory disorder syndrome, sepsis, and septic shock. About 80% of cases were classified as mild or moderate, 13.8% were seriously ill and as many as 6.1% of patients fell into a critical condition. Most patients infected with SARS-CoV-2 show symptoms in the respiratory system such as fever, coughing, sneezing, and shortness of breath [17]. Based on follow-up examinations using the RT PCR method, 75 people (63.56%) were positive for COVID-19.

The screening was carried out using the rapid test method. In order to know the validity of the results, it is necessary to assess the sensitivity and specificity. The current study suggested sensitivity and specificity test for the rapid test tool are 88.66% and a specificity of 90.63 for IgM and IgG [5]. These two parameters’ values were determined using a diagnostic tool that meets the gold standard, namely RT PCR. These two values influence each other. If the sensitivity value increases, the specificity will decrease and vice versa. Based on the calculation results, the sensitivity of the rapid test to the RT PCR results reached 97.33%, and the specificity was 93.02%. The sensitivity of the rapid test resulted in a figure of 97.33% indicating the ability of the test kit to predict a patient with COVID-19, while the specificity of the rapid test resulted in a figure of 93.02% which means that it is good at predicting a patient is not sick with COVID-19. The sensitivity and specificity of an antibody test will vary in results depending on several factors.

Meta-data research conducted in Brazil on 16 rapid test products resulted in a sensitivity and specificity of 82% and 97% [19]. While the results of research conducted in Austria found that the sensitivity and specificity were 98% and 97% [20]. A study conducted in France on 34 positive patients with COVID-19 using three different antibody tests found that all three antibody tests had a sensitivity of around 80% [21]. A study conducted in Germany on 26 samples using four kinds of rapid tests found that the sensitivity ranged from 92.3% - 100% while the specificity ranged from 84%-100% [22]. Research conducted in Finland with 70 samples from COVID patients and 81 control samples found that the sensitivity ranged from 68.3%-97.5% and specificities between 43.8%-81.3% [23]. In Italy, the research was conducted on samples with several times taking, namely seven days, 14 days, and >14 days, the resulting sensitivity was 58.3%, 85.79%, and 100% [24]. Research in Spain on three kinds of rapid test antibodies found sensitivity and specificity values were 100% and 80.6% [25]. A study conducted in China of 150 patients stated that the rapid test antibody has a sensitivity of 71.1% and a specificity of 96.2% [26].

To know the predictive validity, it is known by the positive predictive value (PPV) and the Negative Predictive Value (NPV). PPV is a possible subject that is positively identified by the test equipment.
according to the gold standard. PPV in the screening tool represents the proportion of subjects classified as ill will have a disease in the future [9]. Based on the calculation, the PPV is 96.05%, which means that the proportion of predictions of patients who test positive and will suffer from illness. The use of rapid tests for screening with high sensitivity and PPV values can increase public confidence. However, the predictive value (both PPV and NPV) in the study can not be applied in the population because of differences in the prevalence of cases that occur [9].

There are many rapid antibody test product developments and markets [18], [10]-[30]. Some of these have been evaluated in studies conducted in other countries. However, the sensitivity and specificity values cannot be used as benchmarks in assessing the accuracy of rapid antibody results, but also need to take into account the onset of symptoms and detection of antibodies inhibits sensitivity [24], [31]. Detection of IgG, IgM, and IgA antibodies against SARS-CoV-2 cannot be used as a diagnosis of COVID-19 but as a complement to the RT PCR test in assessing an individual's immune status [24].

This study has several limitations, first: there is no standard for determining the minimum sample size in assessing the sensitivity and specificity of the rapid tests used for screening. Second, no reliable gold standard for serologic assays is currently available for comparative studies, and little literature exists regarding the comparison of the rapid test method for detection of COVID-19. Furthermore, the criteria for assessing the time of disease onset are taken from the last time sample contact with a positive case and may contain imprecision due to subjectivity in the perception of symptoms and timing.

4. CONCLUSION

The sensitivity value of the rapid test antibody used in the study was 97.33% indicating the ability of the test kit to predict a patient with COVID-19. The specific value of the antibody rapid test used in the study was 93.02%, indicating that individuals who are not sick with COVID-19. The positive predictive value (NPP) of antibody rapid test used in the study was 96.05%. The negative predictive value (NPN) of the antibody rapid test used in the study was 95.24%. These results meet the minimum recommendations for the screening method.

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