“It Was Very Comforting to Find Out Right Away.” – Patient Perspectives on Point-of-Care Molecular SARS-CoV-2 Testing in Primary Care

Anni Matthes, Florian Wolf, Jutta Bleidorn, Robby Markwart

1Institute of General Practice and Family Medicine, Jena University Hospital, Friedrich Schiller University, Jena, Thuringia, Germany; 2InfectoGnostics Research Campus Jena, Jena, Thuringia, Germany

Correspondence: Anni Matthes, Institute of General Practice and Family Medicine, Jena University Hospital, Friedrich Schiller University, Bachstr. 18, Jena, Thuringia, 07743, Germany, Tel +49 3641 939 5824, Fax +49 3641 939 5802, Email anni.matthes@med.uni-jena.de

Background: The use of point-of-care tests (POCTs) has been a central strategy to cope with the COVID-19 pandemic. Yet, evidence on the application and consequences of POCTs within medical settings is rare.

Purpose: To assess and understand patient perspectives on molecular point-of-care SARS-CoV-2 testing conducted in primary care.

Methods: We conducted a cross-sectional survey study among patients who were tested with a molecular SARS-CoV-2 rapid test (ID NOW™ COVID-19 rapid test, Abbott) in 13 primary care practices in the state of Thuringia (Germany) from February to April 2021. The following aspects were covered in the questionnaire through rating scales and open text formats: test characteristics, trust in test result, consequences of immediate result, cost amount willing to pay and expectations in the future. Open text answers were categorized; quantitative data were analyzed using descriptive statistics and a Mann–Whitney U-test to reveal differences in cost contribution depending on the test result.

Results: A total of 215 patients from nine family practices and one pediatric practice participated. The immediate availability of the test result was important to the majority of patients (94.3%). 95.7% of patients trusted in their test result. Personal consequences of the immediate test result referred to pandemic measures, certainty of action and reassurance. For further tests, patients were willing to pay between 0€ and 100€ (interquartile range = 10–25€) for the molecular SARS-CoV-2 POCT, regardless of the test result. Expectations of being offered the test again in case of renewed cold symptoms were reported by 96.2%.

Conclusion: Patients highly appreciated molecular SARS-CoV-2 rapid testing conducted in primary care practices. The immediate availability of the test result led to adjustments in patients’ behavior and emotional wellbeing. However, potentially challenging for the implementation of POCTs in primary care practices may be the reimbursement of test costs and patients’ expectations in future situation.

Keywords: POCT, rapid test, COVID-19, acceptance, feasibility

Introduction

Point-of-care tests (POCTs), also known as rapid tests, are in-vitro diagnostic laboratory tests, which are performed in close proximity to the patient and yield test results usually within 30 minutes.1,2 During the COVID-19 pandemic, the public became accustomed to POCTs due to the use of SARS-CoV-2 rapid tests in healthcare facilities and home use. Since April 2021, the use of rapid SARS-CoV-2 antigen tests has been a cornerstone in the German national strategy to cope with the pandemic.3 Besides rapid antigen tests for detecting acute SARS-CoV-2 infections (usually lateral flow assays),4 molecular SARS-CoV-2 POCTs based on nucleic acid amplification techniques with superior diagnostic accuracy have been developed.5,6 However, laboratory testing using the real-time quantitative reverse transcription–polymerase chain reaction (RT-qPCR) technology remains the gold standard in the diagnosis of acute SARS-CoV-2 infection.7 A positive test result has far-reaching consequences for patients and their social network such as quarantine. The rapid availability of the test result could be of particular importance in the context of a disease like COVID-19,
which possesses a significant public health relevance. In addition to the perspectives of medical professionals, the attitude of patients is one important factor in the successful implementation of medical intervention, including diagnostic procedures, in routine healthcare.

In the primary care setting, POCTs should support clinical decision-making, for both patients and physicians. While point-of-care measurement of certain laboratory markers is already established in primary care (e.g., C-reactive protein, D-Dimer, urine dipstick), the development of novel POCTs is omnipresent. Horvath et al. defined key components that should be analyzed in the evaluation of diagnostic tests: analytical and clinical performances, clinical and cost-effectiveness and the broader impact of the test on social, psychological, or organizational consequences (such as patient scheduling, practice management, etc.) in a specific clinical setting. As shown by a review of 40 evaluation reports, the evaluation of novel POCTs targeted for primary care often focused on the diagnostic performance, while evidence gaps exist for the clinical effectiveness and the broader impact of POCTs (e.g., feasibility, acceptance, costs). In order to strengthen the evidence on social, psychological, or organizational consequences of POCTs, it is essential to investigate the perspectives of the actual test users (medical staff) and patients on whom the tests are performed.

The aim of this study was to assess and understand patient perspectives on molecular point-of-care SARS-CoV-2 testing performed in German primary care practices during a COVID-19 infection wave.

**Methods**

**Study Design**

We conducted a descriptive cross-sectional questionnaire-based survey study among patients on whom a molecular SARS-CoV-2 rapid test (ID NOW™ COVID-19 rapid test, Abbott Diagnostics Scarborough, Inc., USA) was performed. The study was conducted in 13 primary care practices in the German federal state of Thuringia from February 15th to April 26th, 2021 when the 7-day incidence ranged between 100 and 200 new SARS-CoV-2 cases per 100,000 inhabitants. In our study, patients included all persons who presented at primary care practices with a suspected SARS-CoV-2 infection. The patients’ perspectives on the following aspects were evaluated: test characteristics, trust in test result, consequences of immediate result, cost amount willing to pay and expectations in the future. The Association of Statutory Health Insurance Physicians Thuringia (The Association of Statutory Health Insurance Physicians Thuringia is a body under public law and represents all outpatient physicians and psychotherapists in the German federal state of Thuringia. It is part of the medical self government and organizes all processes in outpatient medical care of patients insured by the statutory health insurances (Social Security Code (SGB V), § 77)) initiated the study and recruited primary care practices, which were known to be interested in research projects. The analyzer platform and test kits were provided by the manufacturer. By contract between manufacturer and the Association of Statutory Health Insurance Physicians Thuringia, the manufacturer was not involved in study design, implementation and analysis. Evaluation was performed independently without financial remuneration.

During the study period, the use of the molecular SARS-CoV-2 rapid test was restricted to diagnostic testing of patients with typical COVID-19 symptoms (e.g., cough, loss of taste or smell). Physicians had free choice whether SARS-CoV-2 testing was performed and if so, which test was used (laboratory PCR testing or POCT testing). Patients were informed about the nature of the molecular SARS-CoV-2 rapid test by the practice staff. After the test was performed and results were communicated to the patients, they were asked to participate in the study by answering the written questionnaire. The study complies with the declaration of Helsinki. Ethical approval was obtained from the Institutional Research Ethics Board of the Jena University Hospital (Registration No.: 2021–2108-Bef).

**Molecular SARS-CoV-2 Rapid Testing**

The used molecular SARS-CoV-2 rapid test (ID NOW™ COVID-19 rapid test) is based on the isothermal nucleic acid amplification technique for qualitative detection of viruses nucleic acids from nasopharyngeal swabs. According to the manufacturers’ information, the diagnostic sensitivity and specificity are 95.0% and 97.9%, respectively. However, an independent Cochrane review showed that the average sensitivity was 73.0%, while the specificity was 99.7%. As shown by several studies, the sensitivity of the ID NOW™ COVID-19 rapid test is particularly decreased in samples...
with low viral loads, but at high viral loads (corresponding to symptomatic patients), the diagnostic accuracy of the ID NOW™ is comparable to laboratory RT-qPCR testing. In January 2021, local health authorities in the federal state of Thuringia have officially approved the results of the ID NOW™ COVID-19 rapid test as confirmation/non-confirmation of SARS-CoV-2 infections in symptomatic patients without requesting a confirmatory PCR laboratory test. In consequence, patients with a positive ID NOW™ test result were granted the same status and faced equal consequences as patients with a positive laboratory test, such as quarantine orders from their local health authority, certificate of recovery from SARS-CoV-2, and contact tracing.

**Sample**
All patients who underwent molecular SARS-CoV-2 rapid testing in the participating primary care practices were eligible for participation. Practices represented a broad spectrum of the Thuringian practice landscape (small and large as well as rural and urban practices). Eleven family practices and two pediatric practices (including 4 single-handed practices, 6 group practices and 3 outpatient medical care centers) participated in the study and were equipped with the test system.

**Questionnaire**
The questionnaire was designed with the expertise of an interdisciplinary research team consisting of two experienced primary care physicians (J.B., F.W.), a work and organizational psychologist (A.M.) and a biochemist with expertise in primary care laboratory testing (R.M.). To ensure comprehensibility and feasibility as well as to address validity, the questionnaire was piloted by three primary care physicians and five persons without scientific or medical background. The final questionnaire (see Supplementary File 1 for English translation) contained questions regarding patient characteristics (age group, sex, population size of place of residence) and the COVID-19 status (symptom intensity, test result), as well as eight statements on point-of-care SARS-CoV-2 testing to be rated on a five-point Likert scale and three statements to be completed in an open text format. These question formats were used in order to measure the degree of agreement for a series of statements (Likert scale) or to allow a broad range of answers (open text format). The questionnaire consisted of two DIN-A4 pages and could be completed in five to ten minutes. We sent printed questionnaires to the participating practices, with the request to hand them to all eligible patients. Only anonymous data were collected and patients were informed in a written statement that returning the questionnaire to the practice staff implied their consent for anonymous participation in the study.

**Data Analysis**
Raw data from the questionnaire were entered into Microsoft Excel 2010. Statistical analyses were performed using R. All rating items were answered by at least 96.3% of patients. Items in an open text format were answered by 20.0% (Finding out the test result right away, has the following consequences for me …) to 48.4% of patients (I would be willing to pay a maximum of the following amount for the test …). Missing values were excluded from the analysis. We used descriptive statistics to analyse the pattern of patient perspectives. Open text answers were categorized into subcategories in an inductive approach. To investigate the association between the variables test result (grouping variable) and cost amount willing to pay (dependent variable), a Mann–Whitney U-test was performed.

**Results**
In total, 215 patients from nine family practices and one pediatric practice returned completed patient questionnaires. The number of returned questionnaires per practice ranged from 3 to 42 (median = 20). An overview of the characteristics of the participants is provided in Table 1. In our study, 29 patients (15.2%) were tested positive for SARS-CoV-2 by molecular point-of-care testing. The majority of patients (116/198, 58.6%) reported none or mild symptoms, while 76 patients (38.4%) reported moderate or severe symptoms. Most patients (163/204, 79.9%) were between 18 and 65 years old, while 22 participants (10.8%) were older than 65 years. 55.2% of the respondents were female. 60.6% resided in villages or towns with less than 20,000 inhabitants.
General Rating of Molecular Point-of-Care Testing for SARS-CoV-2 in Primary Care Practices

The immediate availability of the test result was of importance for 200 of 212 patients (94.3%) (Figure 1A; see Supplementary File 2 for descriptive results in detail). For 93.4% (197/211) of the participating patients it was important that the rapid test was performed in the primary care practice. A higher reliability compared to other (ie antigen-based) rapid tests was of importance for 89.0% (186/209). 201 of 210 (95.7%) trusted the test result. The great majority (202/212, 95.3%) of patients agreed that they would recommend the molecular point-of-care test for SARS-CoV-2 used in this study to others with symptoms of a cold.

Consequences of Immediate Test Result

For the majority of patients surveyed (208/212, 98.1%) it was of importance to have certainty about potential consequences such as quarantine/self-isolation or informing contact persons immediately (Figure 1B). Forty-three patients provided information on the specific consequence the immediate test result had for them personally. Answers

Table 1 Characteristics of Patients Surveyed

| Number of Patients Surveyed | 215 (100%) |
|----------------------------|-----------|
| Result of molecular SARS-CoV-2 rapid test (n, %*) |          |
| COVID-19 positive | 29 (15.2%) |
| COVID-19 negative | 155 (81.2%) |
| Not specified | 7 (3.7%) |
| NA** | 24 |
| Severity of the symptoms (n, %*) |          |
| None | 30 (15.2%) |
| Mild | 86 (43.4%) |
| Moderate | 58 (29.3%) |
| Severe | 18 (9.1%) |
| Do not know/not specified | 6 (3.0%) |
| NA** | 17 |
| Age group (n, %*) |          |
| 0–17 years | 19 (9.3%) |
| 18–35 years | 49 (24.0%) |
| 36–50 years | 63 (30.9%) |
| 51–65 years | 51 (25.0%) |
| 66–80 years | 20 (9.8%) |
| > 80 years | 2 (1.0%) |
| NA** | 11 |
| Gender (n, %*) |          |
| Female | 111 (55.2%) |
| Male | 90 (44.8%) |
| NA** | 14 |
| Gender ratio (f/m) | 1.23 |
| Population size of place of residence (n, %*) |          |
| < 5000 | 82 (39.9%) |
| 5000–20,000 | 42 (20.7%) |
| 20,000–100,000 | 40 (19.7%) |
| > 100,000 | 32 (15.8%) |
| Do not know/not specified | 8 (3.9%) |
| NA** | 12 |

Notes: *Percentage among valid answers (excluding NA). **No valid answers: item not answered or unclear answer.
were given in short open comments and were clustered in four categories and nine subcategories (Table 2). Statements referred to pandemic-specific aspects such as knowing whether to self-isolate or notifying others. Patients also reported that molecular point-of-care SARS-CoV-2 testing has consequences on their certainty of action either in a general or context-specific (eg work, family) manner: “I can decide not in 2 to 3 days, but immediately, how to behave in my surrounding.” or “I know if my child can go to kindergarten tomorrow – and if I can go to work.” Respondents reported that they had certainty if they could attend COVID-19 vaccination appointments the following day – at the time of the study period (beginning of 2021) vaccination appointments were difficult to get in Germany. Another consequence that patients reported was the reassurance they felt after the test result.

Cost Contribution and Expectations in Similar Future Situations
Among the surveyed patients, 31.9% (66/207) agreed that they would pay the estimated costs of 40€ for the molecular SARS-CoV-2 rapid test, while 37.7% did not agree (78/207). Additionally, patients were asked for the maximum costs they were willing to pay. Answers ranged from 0€ to 100€ (median = 20€, interquartile range = 10–25€, n = 104). A Mann–Whitney test indicated that the amount willing to pay did not differ significantly between patients with a positive (median = 20) and patients with a negative (median = 15) COVID-19 test result ($U = 483.5$, $p = 0.171$). Expectations that a molecular SARS-CoV-2 rapid test will be used in the future if they have similar symptoms were reported by 96.2% (204/212) of patients.
In this study, we surveyed 215 patients on molecular point-of-care SARS-CoV-2 testing performed in primary care practices during a SARS-CoV-2 infection wave in the State of Thuringia, Germany. The rapid test (ID NOW™ COVID-19 rapid test) was approved as equivalent to a PCR laboratory test by local health authorities.

Our study shows that patients accepted and appreciated the characteristics of the molecular SARS-CoV-2 POCT. The vast majority of patients valued the immediate availability of the test result during consultation in primary care practice and showed great confidence in the test result. Having immediate certainty about the potential consequences of the test result, instead of waiting for notification from a central laboratory or public health authority, made a difference for almost all patients. Despite these accepting patient perspectives, our study also identified two aspects that could be challenging for the implementation of a molecular SARS-CoV-2 rapid test in routine primary care: (i) financial aspects and (ii) patient expectations in future situations.

The finding that patients value the immediate test result and the opportunity to discuss implications with their primary care physician within the same consultation has been demonstrated for other indications as well, e.g., lower respiratory tract infection, diabetes, and anticoagulant therapy. Yet, a new and so far COVID-19-specific finding is the difference it made for patients to have immediate certainty about the test result and its consequences. These include self-isolation and informing contact persons, as well as consequences regarding continuation of work and family interaction. With the COVID-19 pandemic being of great public health relevance, consequences of a SARS-CoV-2 test result do not only have an impact on an individual level but also directly affect the society as a whole. Another aspect is the emotional burden that patients face. It has been demonstrated that the waiting period and uncertainty for health-related diagnosis is often provoking anxiety. Patients therefore reported great emotional relief and security after having certainty about their SARS-CoV-2 test result. Thus, the time gain due to the rapid testing had a great impact on patients’ emotional wellbeing as well as their behavioral adjustments.

Considering the implementation of a molecular SARS-CoV-2 rapid test in German routine primary care, the aspect of cost reimbursement may be challenging. In German primary care, relatively few POCT diagnostics are reimbursed and do not routinely include molecular rapid SARS-CoV-2 testing. While expenditures were covered within the study setup, a molecular SARS-CoV-2 test should be reimbursed with 40€ in order to be cost-efficient for the primary care physician. Yet, only 31.9% of patients were willing to cover these costs. Most patients reported that they would pay between 10€ and 25€. It is important to note that in Germany patients are not used to paying for medical services beyond monthly

| Category and Subcategory | Example |
|--------------------------|---------|
| Pandemic measures (n=19) |         |
| Quarantine decision (n=12) | “immediately quarantine”; “not continuing the isolating measures” |
| Notification of contact persons (n=7) | “inform family and friends” |
| Certainty of action (n=20) |         |
| General behavioral certainty (n=8) | “I know how to proceed further:” |
| Work (n=6) | “starting my nursing internship” |
| Family interaction (n=3) | “helps me to decide if I should meet my family” |
| Vaccination appointment (n=2) | “able to attend appointment for corona vaccination” |
| Other (n=1) | “I can buy groceries on the way home from the physician’s office.” |
| Reassurance (n=12) |         |
| Felt secureness (n=6) | “I feel safer, for myself and the people around me.” |
| Emotional relief (n=6) | “mental health”; “relief!!!” |
| No consequences (n=2) | “none” |

Notes: *based on free-text answers of 43 patients, who mentioned a total of 53 personal consequences (some patients stated more than one consequence).
insurance rates. While it remains a challenge for the implementation of molecular SARS-CoV-2 rapid tests, it is still remarkable that participating patients were willing to pay for the test at least to some degree.

At the same time, we anticipate the challenge that POCTs for SARS-CoV-2 might be seen as a new standard and that a rapid test result is expected in similar future medical situations. Even though this aspect has not well been studied so far, Wood et al\textsuperscript{21} conclude for patients from Norway where a POCT is regularly used for managing patients with acute cough that “it appeared to be simply accepted as part of common practice […]” (p.667).

The great acceptance among patients and the rapid detection of SARS-CoV-2 in symptomatic patients advocates for the implementation of molecular SARS-CoV-2 rapid tests in primary care. However, besides costs/reimbursement, many other factors are important for general physicians in the decision to implement a POCT in practice, such as user-friendliness, technical accuracy, frequency of use (disease incidence), clinical guidelines, as well as support, training and quality control.\textsuperscript{29,30} Further studies addressing these aspects are necessary to fully understand the feasibility and utility of molecular SARS-CoV-2 rapid testing in primary care from a clinician’s perspective.

**Strengths and Limitations**
Our study is the first that evaluated patient perspectives on a molecular point-of-care SARS-CoV-2 testing in German primary care. One strength of our study is that the test was evaluated in a clinical routine. While existing evidence and evaluation processes of POCTs often solely focus on their diagnostic performance (sensitivity and specificity), we were able to gain insights into the application and consequences of a SARS-CoV-2 rapid test within the medical setting. Patients answered the written questionnaire immediately after rapid SARS-CoV-2 testing, which minimizes the risk of recall bias.

However, the generalizability of our study is limited to some extent since we only analyzed perspectives of patients in Thuringia. Yet, the included practices represent a certain degree of heterogeneity observed in German primary care because they are located in both rural and urban regions and represent different types of practices. Studies in other countries are necessary to confirm the transferability of our results to other countries. Differences in the healthcare system as well as in culture-specific attitudes toward healthcare should be considered. Additionally, it is likely that patient perspectives on a SARS-CoV-2 test are affected by the pandemic situation and testing regulations at the time. In the beginning of 2021, when our survey took place, only few people (mostly >80 years and/or high-risk populations) in Germany had been vaccinated against COVID-19. Accordingly, the influence of a partial or complete vaccination status on the participants’ perspectives cannot be determined. As of June 2022, 78% of the German population has been vaccinated,\textsuperscript{31} but rigorous testing (incl. rapid tests and laboratory tests) remains a cornerstone of the German COVID-19 strategy. Although there is no evidence that vaccination status affects patient’ attitudes towards molecular SARS-CoV-2 testing in primary care practices, further research on this topic is needed.

**Conclusion**
Based on the perspectives of the surveyed patients, we found a relatively homogenous response pattern for molecular point-of-care SARS-CoV-2 testing in primary care practices: an immediate COVID-19 test result that is equivalent to a PCR laboratory test is of high relevance for patients. It made a difference for patients “to find out right away” and resulted in immediate adjustments of patients’ behavior and emotional wellbeing. Although patients in Germany commonly do not pay for standard healthcare besides insurance fees, patients would, at least partly, pay for a molecular SARS-CoV-2 rapid test. However, the requested cost contribution should not be above 25€. Before implementing molecular SARS-CoV-2 rapid testing in primary care, the question of cost reimbursement as well as perspectives’ of clinicians and clinical stakeholder must be addressed.

**Abbreviations**
POCT, point-of-care test; real-time quantitative reverse transcription–polymerase chain reaction, RT-qPCR; PCR, polymerase chain reaction.
Data Sharing Statement
All data generated or analysed during this study are included in this published article and its supplementary information files (see Supplementary File 3 for Study data).

Ethics Approval and Informed Consent
All methods were carried out in accordance with relevant guidelines and regulations. Informed consent was obtained from all participants. The study protocol including all procedures concerning the study as well as the questionnaire was approved by the institutional research ethics board of the Jena University Hospital (Registration No.: 2021-2108-Bef).

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Author Contributions
All authors made a significant contribution to the work reported, whether that is in the conception, study design, execution, acquisition of data, analysis and interpretation, or in all these areas; took part in drafting, revising or critically reviewing the article; gave final approval of the version to be published; have agreed on the journal to which the article has been submitted; and agree to be accountable for all aspects of the work.

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