Rapid comparative evaluation of SARS-CoV-2 rapid point-of-care antigen tests

Anna Denzler1 · Max L. Jacobs1 · Victoria Witte1 · Paul Schnitzler2 · Claudia M. Denkinger3 · Michael Knop1,4,5

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Abstract

Purpose The objective of this study was to develop a scalable approach for direct comparison of the analytical sensitivities of commercially available SARS-CoV-2 antigen point-of-care tests (AgPOCTs) to rapidly identify poor-performing products.

Methods We present a methodology for quick assessment of the sensitivity of SARS-CoV-2 AgPOCTs suitable for quality evaluation of many different products. We established reference samples with high, medium, and low SARS-CoV-2 viral loads along with a SARS-CoV-2 negative control sample. Test samples were used to semi-quantitatively assess the analytical sensitivities of 32 different commercial AgPOCTs in a head-to-head comparison.

Results Among 32 SARS-CoV-2 AgPOCTs tested, we observe sensitivity differences across a broad range of viral loads (9.8 × 10⁸ to 1.8 × 10⁵ SARS-CoV-2 genome copies per ml). 23 AgPOCTs detected the Ct25 test sample (1.6 × 10⁶ copies/ml), while only five tests detected the Ct28 test sample (1.8 × 10⁵ copies/ml). In the low-range of analytical sensitivity, we found three saliva spit tests only delivering positive results for the Ct21 sample (2.7 × 10⁷ copies/ml). Comparison with published data supports our AgPOCT ranking. Importantly, we identified an AgPOCT widely offered, which did not reliably recognize the sample with the highest viral load (Ct16 test sample with 9.8 × 10⁸ copies/ml) leading to serious doubts about its usefulness in SARS-CoV-2 diagnostics.

Conclusion The results show that the rapid sensitivity assessment procedure presented here provides useful estimations on the analytical sensitivities of 32 AgPOCTs and identified a widely-spread AgPOCT with concerningly low sensitivity.

Keywords Comparison of antigen-detecting rapid diagnostic tests · COVID-19 · SARS-CoV-2 · Test sensitivity · Non-functional AgPOC tests

Introduction

In the SARS-CoV-2 pandemic, lateral flow antigen tests developed as a rapid alternative to SARS-CoV-2 reverse transcriptase quantitative polymerase chain reaction (RT-qPCR)-based diagnostics. Because of their ease of use, lateral flow antigen tests are applicable for point-of-care (POC) as well as self-testing and can, therefore, be incorporated in daily life to support viral containment [1]. In the following, these tests will be referred to as antigen point-of-care tests (AgPOCTs). AgPOCTs are meanwhile widely used for SARS-CoV-2 diagnostics and screening purposes. Currently, several hundred different SARS-CoV-2 AgPOCT brands are commercially available to meet the demand (545 products for professional use are listed by the Federal Institute for Drugs and Medical Devices (Bundesinstitut für Arzneimittel und Medizinprodukte (BfArM)); as of July 27, 2021;
However, the sensitivity and specificity of these are not systematically assessed.

If an AgPOCT is used by a professional operator, it falls under the low-risk category of the European Union directive on In Vitro Diagnostics (IVD), which currently governs marketing authorization for IVD in Europe. Under this directive, manufacturers can still self-certify SARS-CoV-2 AgPOCTs and waive independent quality control before these products enter the market. The validation of available AgPOCTs is, therefore, not assured in the view of the Paul Ehrlich Institute (PEI), Federal Institute for Vaccines and Biomedical Products, Germany. Furthermore, the PEI reports on evidence of counterfeiting. A new regulation governing independent validation by specialized and certified reference laboratories is planned, but will only become effective in May 2022 at the earliest (https://www.pei.de/DE/newsroom/hp-meldungen/2020/200323-covid-19-nat-tests.html;jsessionid=F786872E8B85959A8E8DA2B8FCB3ABE00.intranet222?nn=169730).

If an AgPOCT is distributed for layperson use, it falls under a higher-risk category and requires independent validation. This validation of sensitivity is currently performed by the PEI together with reference laboratories and a list with examined AgPOCTs passing their criteria is provided [3]. AgPOCTs failing the comparative evaluation by PEI will be removed from the list provided by the BfArM [2]. This list, however, comprises only products from manufacturers or distributors, which also registered their products for listing (https://www.bfarm.de/DE/Medizinprodukte/Aufgaben/Spezialthemen/Antigentests/_node.html), rendering the absence of an AgPOCT from this list difficult to interpret.

Many in-depth AgPOCT characterization studies show that AgPOCT sensitivities can vary substantially. One study reporting on the validation of 122 AgPOCs was recently published [4]. The authors found that 26 AgPOCTs do not fulfill the required minimum sensitivity, clearly illustrating that many circulating AgPOCTs are insufficiently sensitive. In addition to this, significant brand-to-brand and lot-to-lot variations were observed [5]. These circumstances urge the need for an easy-to-use method to quickly assess AgPOCTs at market entry and periodically thereafter for post-implementation quality control.

In this study, we sought to establish a procedure to rapidly evaluate the sensitivity of a large number of AgPOCTs using a small test sample panel and several tests per product. To this end, we developed a strategy involving pooled samples and four different dilution steps from high to low viral loads and generated several hundred aliquots thereof. Using this approach we then investigated 32 AgPOCTs, which included mainly tests currently in use in the local area (Heidelberg, Germany). We compared the results with data from the literature, which enabled us to conclude the validity of our approach and the performance of the products investigated.

**Methods**

**Study design**

We tested the analytical sensitivity of a large number of commercially available AgPOCTs by applying pooled samples from nasopharyngeal swabs with defined SARS-CoV-2 viral loads (given in cycle threshold (Ct)) including Ct16, Ct21, Ct25, and Ct28 (~9.8×10^8 to ~1.8×10^5 genome copies per ml) as well as a pooled sample obtained from SARS-CoV-2 negative tested persons. Pools were generated using anonymized remnant swab sample material collected for the clinical diagnosis of a SARS-CoV-2 infection by RT-qPCR carried out by the Center for Infectious Diseases, Virology, Heidelberg University Hospital, Germany. Pharyngeal swab specimens were collected through the nose (nasopharyngeal) and contained in viral transport medium (VTM). Per test, 50 µl of the samples were mixed with the provided lysis buffer of each AgPOCT, and the tests were performed strictly according to the manufacturer’s instructions. After the recommended incubation time, we acquired images of the test chambers using a Panasonic Lumix DMC-G70 camera equipped with a Panasonic H-FS12060 objective. We tested AgPOCTs at least in duplicates with the corresponding test samples. We quantified test results by measuring the background-corrected signal intensities of the test (T) band versus control (C) band in ImageJ (v1.53c) using the Gels analysis function usually used for quantification of Western Blot bands. For qualitative evaluation of the visibility of the test bands (positive versus negative score), RGB pictures of AgPOCT results from randomly chosen replicates were evaluated independently by three individuals in a blinded manner. Furthermore, all test results were scored independently by another person.

**Preparation of test samples from nasopharyngeal swabs**

Anonymized, remnant nasopharyngeal swab samples tested positive and negative for SARS-CoV-2 were obtained between May and July 2021 from the Center for Infectious Diseases, Virology, Heidelberg University Hospital, Germany. Samples were stored in VTM. The Ct16, Ct21, and negative test samples were prepared by pooling of 12–15 nasopharyngeal swab samples. Cell debris and other solids were removed by centrifugation at 400g for 10 min and subsequent transfer of the supernatant.
into a new tube. Viral RNA was isolated from pools by manual lysis and automated RNA extraction using the QIAamp Viral RNA Mini kit (Qiagen) on a QIAcube Connect device (Qiagen). Ct values of sample pools were determined by RT-qPCR analysis using the LightMix® Modular Sarbecovirus SARS-CoV-2 (TIB Molbiol) with the LightCycler® Multiplex RNA Virus Master (Roche) and LightCycler480 II (Roche). Subsequently, pools were supplemented with 2% TritonX-100 and complete Ultra protease inhibitor (Roche) and if needed adjusted with dilution buffer [2 mg/ml BSA, 0.9% NaCl, protease inhibitor]. Ct25 and Ct28 test samples were prepared by dilution of the Ct21 test sample in dilution buffer. Samples were aliquoted (120 µl), immediately frozen on dry ice, and stored at -80 °C. Viral loads of Ct test samples were estimated based on a standard curve determined using the quantitative INSTAND SARS-CoV-2 reference samples 1 and 2 (Ch07469 and CH07470 with defined viral loads of 10,000,000 and 1,000,000 copies/ml; Supplemental Figure S6). For AgPOCT testing, samples were freshly thawed on ice before use. Each time, test samples were validated using a LumiraDx SARS-CoV-2 microfluidics POCT system [6].

**AgPOCTs evaluated in this study**

We included a total of 32 AgPOCTs available at local supermarkets, pharmacies, and drugstores as well as on several online trade platforms (Table 1). Specific AgPOCTs will be referred to as the respective manufacturer’s name (in italic in Table 1). The inspected AgPOCTs include both, tests for professional in vitro diagnostics use (#1–14) as well as tests temporarily licensed for self-testing in Germany (#15–32) by the Federal Institute for Drugs and Medical Devices [2] (Supplemental Figure S5). The majority of AgPOCTs available were nasal or nasopharyngeal swab tests except BTNX, Ritter, Joinstar, Realy (#11–14) among the tests for professional in vitro diagnostics use (#1–14) and Hygisun, fameditec (#30–32) among the self tests, which are all saliva spit tests, as well as Watmind (#29), which is a saliva swab test.

For Lepu medical (#20 in Table 1), the AgPOCT with the poorest results in our study, we purchased different versions and additional batches for in-depth characterization (Table 2).

**Results**

**Generation of test samples for standardized AgPOCT evaluation**

In the present study, we sought to establish a standardized procedure to rapidly assess the sensitivities of a large number of SARS-COV-2 AgPOCTs. To this end, we generated a collection of test samples from pooled nasopharyngeal swabs from SARS-CoV-2 positive-tested and negative-tested individuals. Ct values of the SARS-CoV-2 positive pools were determined by RT-qPCR and test samples were prepared accordingly. The test sample collection comprised four SARS-CoV-2 positive pools with defined viral loads (Ct16, Ct21, Ct25, and Ct28) and one SARS-CoV-2 negative pool. The positive pools covered high (Ct values < Ct24) and moderate (Ct values between Ct24 and Ct30) loads of SARS-CoV-2 genetic material. Per test sample, >200 aliquots with 120 µl sample volume each were prepared, allowing a quick and standardized evaluation of the analytical sensitivities of a large number of different AgPOCTs.

We estimated that our test sample collection covers a range from $9.8 \times 10^8$ (Ct16) to $1.8 \times 10^5$ (Ct28) SARS-CoV-2 genome copies per ml (Supplemental Figure S6). We qualitatively validated the test sample collection using a LumiraDx Covid-19 antigen test device. This microfluiddic POCT system was manufactured in Europe and able to detect the viral antigens in the most diluted sample. This is consistent with previous reports on the high sensitivity of the LumiraDx device [6]. We used 50 µl of a test sample, each for the LumiraDx analysis and for all AgPOCTs evaluated in this study, as described before [7, 8]. All four SARS-CoV-2 positive test samples tested positive for SARS-CoV-2, while the negative test sample was recognized as negative in the LumiraDx analysis.

**Quantitative and qualitative assessment of AgPOCT analytical sensitivity**

We tested a total of 32 AgPOCTs (Table 1). We purchased 12 AgPOCTs from local resellers (pharmacies, drugstores, supermarkets) and 20 products online. We performed the tests over 10 days, with the help of four students, during the course of four weeks. For each product, we used freshly thawed aliquots of the Ct21, Ct25, and Ct28 test samples as well as the negative sample. We conducted two to four replicates per product and acquired images of each of the tests at the time points specified by the manufacturers. The Ct16 test sample was only used for AgPOCTs that had a low performance with the Ct21 test sample. For quantitative evaluation, signal intensities of the test ($T$) and the control ($C$) bands were measured and the ratio of these values ($T/C$ ratio) was determined (Fig. 1a). In addition, we scored a binary (positive or negative) test result using visual inspection of the images by four different persons (Fig. 1b).

For 31 of 32 investigated AgPOCTs, an average $T/C_{C_{25}} > 0$ was determined for all positive test samples and not for the negative control sample (Fig. 1a). This indicates that the digital quantification detects test band signals for 31 AgPOCTs using the Ct25 test sample, albeit some test bands
| #  | Supplier                                      | AgPOCT name                                      | Specifications                              | Sample type       | Use          | Distributor       |
|----|----------------------------------------------|--------------------------------------------------|---------------------------------------------|-------------------|--------------|-------------------|
| 1  | Abbott Rapid Diagnostics Jena GmbH           | Panbio COVID-19 Ag Rapid test device (nasal)     | REF: 41FK11                                 | Nasal swab        | pro Online trade |
|    |                                              |                                                  | LOT: 41ADG244A                              |                   |              |                   |
| 2  | Healgen Scientific Limited Liability Company | Coronavirus Ag Rapid Test Cassette (Swab)        | REF: GCCOV-502a                              | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: 2012650                                |                   |              |                   |
| 3  | RapiGEN, INC                                | Biocredit COVID-19 Ag—one step Rapid Test       | REF: G61RHA20                                | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: H0730975D                               |                   |              |                   |
| 4  | Beijing Beier Bioengineering Co., Ltd        | COVID-19 Antigen Rapid Test Kit                  | REF: not specified                          | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: 20210201                               |                   |              |                   |
| 5  | möLab GmbH                                  | mō-screen Testkit Corona Antigen                | REF: 0230005B1                               | Nasal/nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOS: 2104072                                |                   |              |                   |
| 6  | Biomerica, Inc                              | COVID-19 Antigen Rapid Test                     | REF: 1509A-251                               | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: COV6686                                 |                   |              |                   |
| 7  | Joysbio (Tianjin) Biotechnology Co., Ltd     | SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) | REF: G10313                                  | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: 2021011607                              |                   |              |                   |
| 8  | Safecare Biotech (Hangzhou) Co., Ltd        | COVID-19 Antigen Rapid Test Kit (Swab)           | REF: COV Ag-6012                             | Nasopharyngeal swab | pro Online trade |
|    |                                              |                                                  | LOT: COV21040606                             |                   |              |                   |
| 9  | Hangzhou Testsea Biotechnology Co., Ltd     | Testsealabs COVID-19 Antigen Test Cassette Corona Antigen | REF: 2020013 vB/10 LOT: TL1C05             | Nasal swab        | pro Online trade |
|    |                                              |                                                  |                                             |                   |              |                   |
| 10 | ACON Biotech (Hangzhou) Co., Ltd            | Flowflex SARS-CoV-2 Antigen-Schnelltest (Selbsttest) | REF: L031-11855                             | Nasal swab        | pro Online trade |
|    |                                              |                                                  | LOT: COV1030052                              |                   |              |                   |
| 11 | BTNX Inc                                    | Rapid Response COVID-19 Antigen Rapid Test Cassette | REF: COV-2C25B                               | Saliva (spit)     | pro Online trade |
|    |                                              |                                                  | LOT: COVG21030089                            |                   |              |                   |
| 12 | Joysbio (Tianjin) Biotechnology Co., Ltd/Litt/Ritter | Easy Check Spittest SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold) | REF: COV-AG-20/G10313 | Saliva (spit)     | pro Online trade |
|    |                                              |                                                  | LOT: 20210202                                |                   |              |                   |
| 13 | Joinstar Biomedical Technology Co., Ltd     | COVID-19 Antigen Rapid Test (Latex)              | REF: RPBH12360                               | Saliva/sputum (spit), stool | pro Online trade |
|    |                                              |                                                  | LOT: COV2103002L                             |                   |              |                   |
| 14 | Hangzhou Realy Tech Co., Ltd               | Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Device (Saliva) | REF: K590516D | Saliva (spit) | pro Online trade |
|    |                                              |                                                  | LOT: 202101022                               |                   |              |                   |
| 15 | nal von minden GmbH                         | NADAL COVID-19 Ag Test                           | REF: 243103 N-20H                            | Nasal swab        | lay Online trade |
|    |                                              |                                                  | LOT: 175363                                  |                   |              |                   |
|    |                                              |                                                  | BfArM GZ: 5640-S-045/21                     |                   |              |                   |
| 16 | SD Biosensor                                | SARS-CoV-2 Rapid Antigen Test                    | REF: 9901-NCOV-01G                           | Nasal swab        | lay Online trade |
|    |                                              |                                                  | LOT: QCO390092I                              |                   |              |                   |
|    |                                              |                                                  | BfArM GZ: 5640-S-025/21                     |                   |              |                   |
| 17 | Beijing Hotgen Biotech Co., Ltd             | Coronavirus (2019-nCoV)-Antigenstest             | REF: 426020532859                           | Nasal swab        | lay Supermarket Pharmacy |
|    |                                              |                                                  | LOT: W2021032500/602/1500                   |                   |              |                   |
|    |                                              |                                                  | BfArM GZ: 5640-S-057/21                     |                   |              |                   |
| 18 | Guangzhou Wondfo Biotech Co., Ltd           | 2019-nCoV Antigen Test (Lateral Flow Method)     | REF: W634P0021                              | Nasal swab        | lay Supermarket |
|    |                                              |                                                  | LOT: W634014116                             |                   |              |                   |
| 19 | Teda Laukoetter Technology GmbH             | COVID-19 Antigen Schnelltest (kolloidales Gold) ANBIO Corona Antigen Nasentupfer | REF: A6061214 | Nasal swab     | lay Drug store |
|    |                                              |                                                  | LOT: 2021046133/461310/036138               |                   |              |                   |
|    |                                              |                                                  | BfArM GZ: 5640-S-079/21                     |                   |              |                   |
had extremely weak signal intensities (Fig. 2). For almost all AgPOCTs, no signal could be measured when the SARS-CoV-2 negative test sample was applied. Only for Jointstar, one replicate of the negative test sample resulted in a false positive test band indicated by a $T/C_{\text{Neg.}} > 0$. In contrast to the more sensitive digital quantification, the visual inspection did only score a positive result for 28 of 31 AgPOCTs with a $T/C_{C_{25}} > 0$ (Fig. 1b, Fig. 2). This is also true for the visual assessment of the results of technical replicates. For example, for Jointstar, the negative sample with a $T/C_{\text{Neg.}} > 0$ scored negative in the visual inspection. We could not establish a specific $T/C$ value threshold to explain the results of the visual assessment, indicating that these ratios are product-specific. This can be explained by different dyes, and by the fact that the visual assessment was conducted using color vision, while for the $T/C$ quantification grayscale images
Table 2  Lepu medical AgPOCT products investigated in this study

| Product name | Packaging size (use) | Specifications | Production and use-by date |
|--------------|----------------------|----------------|----------------------------|
| SARS-CoV-2 Antigen Rapid Test Kit (Colloidal Gold Immunochromatography) CE | Pack of 25 tests (pro) | Barcode: 6921807601020<br>LOT: 21CG2713X<br>Barcode: same as above<br>LOT: 21CG2715X | Production date: 06/03/2021<br>Use-by date: 06/03/2022 |
| NASOCHECKcomfort SARS-CoV-2 Antigen Schnelltest Immunchromatographischer Test (kolloidales Gold). Schnell & angenehm! Einfache Anwendung im vorderen Nasenbereich | Pack of 25 tests (lay) | REF: CG2725(N)<br>Barcode: 4260716970042<br>LOT: 21CG2722X<br>BfArM GZ: 5640-S-104/21 | Production date: 31/03/2021<br>Use-by date: 31/03/2022 |
| NASOCHECKcomfort SARS-CoV-2 Antigen-Schnelltest. Zur Eigenanwendung. Schnell & angenehm! Einfache Anwendung im vorderen Nasenbereich | Single pack (narrow) (lay) | REF: CG2701N<br>Barcode: 4260716970059<br>LOT: 21CG2727X<br>BfArM GZ: 5640-S-104/21 | Production date: 11/04/2021<br>Use-by date: 11/04/2022 |
| NASOCHECKcomfort SARS-CoV-2 Antigen-Schnelltest. Zur Eigenanwendung. Schnell & angenehm! Einfache Anwendung im vorderen Nasenbereich | Single pack (thick) (lay) | REF: CG2701N<br>Barcode: 4260716970059<br>LOT: 21CG2720X<br>BfArM GZ: 5640-S-104/21<br>REF, barcode, BfArM GZ: same as above<br>LOT: 21CG2724X | Production date: 04/04/2021<br>Use-by date: 04/04/2022 |

AgPOCT products made by Beijing Lepu Medical Technology Co., Ltd. are referred to as Lepu medical AgPOCTs. Lepu medical AgPOCTs were purchased on different online trade platforms. For each Lepu medical AgPOCT, product name, packaging size, and intended use (professional (pro) versus layman (lay) use), reference/barcode number, LOT number as well as BfArM GZ number if applicable are indicated. Production and use-by date are noted in the last column.
were used. Furthermore, we observed large coefficients of variation (CV) for some of the tests, in particular for samples with very small T/C ratios, emphasizing weak signals close to the detection limit of the digital quantification (Supplemental Figure S1c).

We grouped the tested AgPOCTs into categories with low (Group III), medium (Group II), and high (Group I) sensitivity based on the reliability to detect a given SARS-CoV-2 positive sample. A sample was considered reliably detected by a given AgPOCT when all or the majority of replicates (at least two out of three or three out of four replicates) of a given sample were scored positive. If none or the minority of replicates of a given sample was detected by the corresponding AgPOCT, reliability requirements were not met.

One exception was Lepu medical (Table 1, AgPOCT #20; Table 2, last row), which did not fulfill the requirements for any of these groups. For Group III AgPOCTs with the lowest sensitivity, the minimum criterion was the reliable detection of the Ct21 sample. Lepu medical tests failed this, as these did not even reliably score a positive result with the Ct16 sample (9.8 × 10^8 copies/ml; Fig. 1b, Fig. 2). To investigate this product further, we used individual unprocessed nasal/nasopharyngeal swab samples with low Ct values (Ct13.3 to Ct18.4) on this and another poor performing product. Comparison of these results to the Ct16 test sample confirmed the low sensitivity of the Lepu medical test (Supplemental Figure S2). Only for samples with very low Ct values (Ct~13), T/C ratios were obtained that can be detected easily visually (Supplemental Figure S2a). This suggests that this product is not completely non-functional, but largely insensitive. Since Lepu medical AgPOCTs have been widely used in Germany and other European countries we retrieved several Lepu medical products available on different online trade platforms (Table 2). These included two different batches of a Lepu medical product intended for professional use (Table 2, first row, no BfArM GZ number, CE mark). Additionally, we included different batches and deviations of the Lepu medical AgPOCT described before (Table 1, AgPOCT #20; Table 2, last three rows). These products were provided with the same BfArM GZ number (5640-S-104/21), which cannot be found on the BfArM list anymore (Supplemental Figure S5). We investigated their performance in direct comparison in multiple replicates using Ct16, Ct21, and Ct25 test samples (Supplemental Figure S3). This revealed high variation of the determined T/C ratios, with coefficients of variation (CV) ranging from 0.26 to 1.54 and a median CV of 0.50 (Supplemental Figure S4c). In contrast, the median CV of the other 31 investigated AgPOCTs with Ct16-25 test samples was 0.11 (Supplemental Figure S1c). This indicates a larger fluctuation of the test results not only for different implementations of the Lepu medical AgPOCTs but also for different batches of the same Lepu medical product, compared to all other AgPOCTs investigated in this study.

AgPOCTs in Group III only reliably detected the Ct21 sample (2.7 × 10^7 copies/ml) and included Hygisun, Joinstar, and Ritter. Of note is that all of them are saliva-based spit tests (Table 1), which are provided with a considerably larger amount of lysis buffer (500–1000 µl lysis buffer; Fig. 1c) than most other AgPOCTs. This results in an increased dilution of the test sample compared to AgPOCTs for nasal samples, which are provided on average with 320 µl lysis buffer (Fig. 1c). The higher dilution of the sample together with the possibility of lower virus concentration in saliva versus nasal or nasopharyngeal swabs may further influence the analytical sensitivity of these AgPOCTs.

The large majority of the investigated AgPOCTs (23 out of 32) delivered visible positive results with the Ct25 sample (1.6 × 10^6 copies/ml, Group II). Among these 23 AgPOCTs, positive scoring was fully reproducible in all replicates for 17 AgPOCTs. AgPOCTs intended for professional use (sorted ascending according to T/C_{Ct25}: Safecare, Realy, Healgene, ACON, Beier, Testsea, BTNX, and Biomerica) largely cluster in the upper half of the T/C_{Ct25} ranking, while tests licensed for self-testing largely cluster in the lower half (sorted ascending according to T/C_{Ct25}: Sanicom, fameditec, OFM, Deepblue, NanoRepro, nal von minden, Teda, Laihe and Boson). Interestingly, among both tests for professional and for layman use, saliva spit tests (Realy, Sanicom, fameditec) appear largely inferior compared to nasal swab tests in this setting except for of BTNX, which is the sixth-highest ranked AgPOCT among all investigated tests. Six AgPOCTs in Group II (sorted ascending according to T/C_{Ct25}: Sanicom, fameditec, OTM, Deepblue, SD Biosensor, Abbott, and Wondfo) failed in one out of three to four replicates to detect the Ct25 sample, which is represented by larger CV values ranging from 0.26 to 0.87 (Supplemental Figure S1c).

Using the Ct28 test sample (1.8 × 10^5 copies/ml), 14 out of 32 AgPOCTs yielded a T/C_{Ct28} > 0; however, only five reliably scored a positive result in the visual investigation (Group I). These include (sorted in ascending order according to T/C_{Ct28}) möLab, Medice, MP, Clongene, and Watmind, three of which are temporarily licensed for self-testing (Table 1, Supplemental Figure S5). All, except möLab, delivered a positive visual result in all three replicates.

Taken together, the data presented here demonstrate that the different SARS-CoV-2 AgPOCTs available deviate largely in the analytical sensitivity of the lateral flow test stripes and provided buffer systems, corresponding more than two orders of magnitude of viral genome copies per ml (9.8 × 10^8 to 1.8 × 10^5). Additionally, we revealed great variation in results delivered with different Lepu medical AgPOCT versions and batches emphasizing the need for regular quality monitoring.
Currently, there are more than 500 different products available for SARS-CoV-2 diagnostics, many of which lack an independent assessment of their performance. In most cases, the clinical sensitivity values provided by the manufacturer (examples in Fig. 1c) are far > 90%. However, detailed information on specimen collection and viral loads are usually not provided rendering these values largely inconclusive and misleading for laymen. Considering that individual products use different antibodies in varying amounts with different specificities and affinities sometimes recognizing different...
proteins in the viral particle with unequal abundances, and diverse staining methods, these conspicuously similar values for clinical sensitivity given by the manufacturers are also unlikely. Therefore, an independent, rapid and critical evaluation of AgPOCTs available is required to determine the realistic performance of AgPOCT relevant to the daily user and especially to identify poor performing products. Given the huge number of products available for rapid SARS-CoV-2 diagnostics, in-depth studies evaluating the quality of AgPOCTs in a time-intensive procedure will not be available any time soon for all products available.

We developed a straightforward strategy to quickly evaluate the technical sensitivity of AgPOCTs for SARS-CoV-2. We generated four SARS-CoV-2 positive reference samples by pooling several individual samples with high viral loads followed by RT-qPCR analysis and subsequent adjustment of the pools to the desired Ct values with buffer. Reference samples with higher Ct values were generated by further dilution of the above-mentioned pools. This approach results in mixed samples and the impact of contaminations in individual samples needs to be considered: potentially, inhibitory contaminations or contaminations that are causing false-positive results may be present in one or the other sample used for the pools. However, such samples are usually an exception. Furthermore, the pooling and Ct adjustment procedure results in a 15- to 20-fold dilution of such constituents so that their impact is reduced if not completely marginalized. Nevertheless, one needs to consider that individual AGPOCTs may be influence by such contaminations. The reference samples were adjusted to span the relevant dynamic range of the typical sensitivity of AgPOCTs ranging from Ct28 to Ct16 (corresponding to approximately $1.8 \times 10^7$ to $9.8 \times 10^8$ SARS-CoV-2 genome copies per ml). Lower Ct values correspond to higher viral loads. The Ct value alone cannot inform about the fraction of viable virus; however, multiple studies showed that high viral loads (Ct values < Ct24) upon detection of a SARS-CoV-2 infection relate to higher morbidity and infectivity potential (summarized in [9]). Therefore, AgPOCTs are, despite their inferior sensitivity compared to laborious RT-qPCR-based SARS-CoV-2 diagnostics, a critical tool in aiding disease prognosis and viral containment by detecting these high to moderate viral loads.

Using these reference samples, we were able to group 32 commercially available products into AgPOCT groups with high, average and low sensitivity (Group I-III). Most importantly, we identified one product that did not detect any of the test samples and, therefore, is considered not suitable for SARS-CoV-2 diagnostics.

The majority of tests investigated in this study reliably detected the Ct25 test sample as SARS-CoV-2 positive (Group II). Some of these AgPOCTs have been thoroughly characterized, including Abbott, RapiGEN, Healgen, nal von minden, and SD Biosensor by Corman et al. [7] and others [4, 8–23]. Corman et al. determined 95% limits of detection for each AgPOCT using 138 SARS-CoV-2 positive clinical samples with viral loads ranging from $1.9 \times 10^4$ to $2.8 \times 10^9$ genome copies per ml. Among the AgPOCTs also tested in this study, Healgen was most sensitive, followed by Abbott, SD Biosensor, and nal von minden—all with a 95% limit of detection between $2.3$ and $9.3 \times 10^6$ SARS-CoV-2 genomes per swab. In contrast, for RapiGEN, a 95% limit of detection more than three orders of magnitudes lower was found. This discrepancy in performance between RapiGEN and the above-mentioned products is supported by other studies [24]. In our analysis, this trend is reflected as well even though we cannot resolve the limits of detection in such great detail: For Healgen and nal von minden, detection of the Ct25 test sample ($1.6 \times 10^6$ copies/ml) was robust with all replicates being positively scored. For RapiGEN, Ct25 test sample detection was less reliable, and based on the $TIC_{Ct25}$ this product is ranked in the lowest quarter among all AgPOCTs investigated.

Among the 32 investigated AgPOCTs, we identified four reliably well-performing AgPOCTs, which detected the Ct28 test sample ($1.8 \times 10^5$ copies/ml) as SARS-CoV-2 positive in all replicates (Group I). Amongst these, Watmind was ranked highest in our evaluation. This AgPOCT is also among the best-performing three AgPOCTs out of 122 tested products with a sensitivity of 82% in samples with Ct values ranging from Ct17 to Ct35 corresponding to viral loads of $>10^8$ to $10^9$ SARS-CoV-2 genome copies per ml [4].

Group III includes AgPOCTs with lower analytical sensitivity as these only detected the Ct21 test sample as SARS-CoV-2 positive. For Joinstar, using Latex beads for visualization, evidence provided by Scheiblauer and colleagues [4] suggests that this test is non-functional with 0% sensitivity.
for all sample panels supporting the low ranking of Joinstar in this study. In our analysis, we detected weak bands for the Ct21 and Ct16 test samples; however, these were considerably weaker than for all other tests suggesting the possibility that latex beads used for visualization do fail to produce a strong signal. Besides Joinstar, Ritter and Hygisun, both saliva spit tests as well, showed low sensitivities in our studies. While we could not find independent evaluation studies for these products, both can be found on the BfArM list (as of July 23, 2021; Supplemental Figure S5).

Fig. 2 Representative images of SARS-CoV-2 AgPOCTs lateral flow test stripes treated with corresponding Ct test samples. Contrast settings were optimized for each AgPOCT example image set to ensure the best visibility of the test bands. AgPOCT example images are arranged (from top left to bottom right) according to the ranking presented in Fig. 1. The red line indicates the limit of reliable detection (see Fig. 1a, b). Arrowheads highlight positions of control (C) and test (T) bands.
Among the low ranked AgPOCTs, the sensitivity of the Lepu medical AgPOCT was exceptionally low as this test failed to deliver a visible positive test result in most replicates, even for the Ct16 sample. In addition to its poor performance in SARS-CoV-2 diagnostics, out of 20 performed Lepu medical tests, three tests technically failed, indicated by the absence of the control band. Importantly, this AgPOCT is a popular product in the area where this study was conducted and is still available at many drugstores and supermarket chains. Of note is that Lepu medical differs from other AgPOCTs in its design and sample application. Technical failure did not occur in any of the other AgPOCTs, in which the immunochromatography paper is embedded in the common plastic cassette. In other studies, Lepu medical AgPOCT products performed better [4, 25], e.g. in the setting of Scheiblauer et al. [4], a sensitivity of 100% was found for a Lepu medical AgPOCT and test panel members with Ct values ranging from Ct17 to Ct25. As the AgPOCTs used in these studies are not specified with reference/product and LOT number, it is possible that a different Lepu medical product or batch was used. We purchased different Lepu medical AgPOCT products available online and compared performances on the pooled test samples and raw, unprocessed swab samples (Supplemental Figure 3). We tested two batches of a CE-marked product and four Lepu medical with the same BfArM GZ number, but different packaging versions (Table 2). Indeed, we identified two Lepu medical products performing better than the Lepu medical AgPOCT and test panel members with Ct values ranging from Ct17 to Ct25. As the AgPOCTs used in these studies are not specified with reference/product and LOT number, it is possible that a different Lepu medical product or batch was used. We purchased different Lepu medical AgPOCT products available online and compared performances on the pooled test samples and raw, unprocessed swab samples (Supplemental Figure 3). We tested two batches of a CE-marked product and four Lepu medical with the same BfArM GZ number, but different packaging versions (Table 2). Indeed, we identified two Lepu medical products performing better than shown in Fig. 1; however, these performances were not reproducible with other batches of the same product (Supplemental Figure 3), indicating batch-specific variation of the quality. Issues regarding the quality of rapid AgPOCTs are reported not only for SARS-CoV-2 [1], but also other infectious diseases such as malaria [26]. This, again, emphasizes the importance of a simple method to assay the performance of an AgPOCT product and corresponding batches. In summary, a comparison with the same strategy constitutes a viable solution to rapidly assess the sensitivity of SARS-CoV-2 AgPOCTs.

It is important to mention that the sensitivity of an AgPOCT is the product of multiple factors; the sensitivity of the test stripe and buffer system used are important contributing factors, but not the only ones. Another is the volume of the lysis buffer provided with each AgPOCT. The volume varies between tests of different manufacturers resulting in a 2.6- to 20-fold dilution of the samples (Fig. 1c). In this study, we did not correct the test results for these different dilution factors, because the sample dilution is an internal property of each AgPOCT. By using the same sample volume for each AgPOCT we also neglect potential differences in swab properties, such as absorption volume or sample specimen (saliva, nasal or nasopharyngeal samples), which affect the diagnostic sensitivity of AgPOCTs. However, we note that for tests based on nasal swabs the used volume of 50 μl approximates the quantity absorbed by these swabs [7]. Furthermore, the AgPOCT-specific instructions for self-sampling, which will influence how carefully a sample is collected, can also influence the diagnostic sensitivity of a test. In light of these considerations, we want to emphasize that this evaluation method only and exclusively focuses on comparing the technical sensitivity of the lateral flow test strips from different test manufacturers, in combination with the provided lysis buffers.

The procedure presented here involving a reduced test sample collection and minimal labor represents a feasible strategy for prompt evaluation of available AgPOCTs for their usability in SARS-CoV-2 diagnostics. We provide a useful estimation of the limits of detection for the investigated AgPOCTs as the dimensions and trends are comparable to results from much more laborious in-depth studies. Importantly, using this approach, we revealed very heterogeneous results for the Lepu medical AgPOCT, which precludes in our opinion the use of this product (or product family) for SARS-CoV-2 diagnostics. In conclusion, we suggest this procedure as a rapid alternative to investigate COVID-19 AgPOCTs in the absence of reliable data that validate the performance of a specific product and related batches.

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Declarations

Conflict of interests The authors have declared no competing interests relevant to this article.

Ethics approval Samples used in this study were access material from nasopharyngeal swabs derived from individuals, which were tested for a SARS-CoV-2 infection by RT-qPCR carried out by the Center for Infectious Diseases, Virology, Heidelberg University Hospital. Swab samples were anonymized at any time and exclusively used for SARS-CoV-2 detection using AgPOCTs in this study.

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