Performance of Saliva, Oropharyngeal Swabs, and Nasal Swabs for SARS-CoV-2 Molecular Detection: a Systematic Review and Meta-analysis

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ABSTRACT
Nasopharyngeal (NP) swabs are considered the highest-yield sample for diagnostic testing for respiratory viruses, including SARS-CoV-2. The need to increase capacity for SARS-CoV-2 testing in a variety of settings, combined with shortages of sample collection supplies, have motivated a search for alternative sample types with high sensitivity. We systematically reviewed the literature to understand the performance of alternative sample types compared to NP swabs. We systematically searched PubMed, Google Scholar, medRxiv, and bioRxiv (last retrieval 1 October 2020) for comparative studies of alternative specimen types (saliva, oropharyngeal [OP], and nasal [NS] swabs) versus NP swabs for SARS-CoV-2 diagnosis using nucleic acid amplification testing (NAAT). A logistic-normal random-effects meta-analysis was performed to calculate % positive alternative-specimen, % positive NP, and % dual positives overall and in subgroups. The QUADAS 2 tool was used to assess bias. From 1,253 unique citations, we identified 25 saliva, 11 NS, 6 OP, and 4 OP/NS studies meeting inclusion criteria. Three specimen types captured lower % positives (NS [82%, 95% CI: 73 to 90%], OP [84%, 95% CI: 57 to 100%], and saliva [88%, 95% CI: 81 to 93%]) than NP swabs, while combined OP/NS matched NP performance (97%, 95% CI: 90 to 100%). Absence of RNA extraction (saliva) and utilization of a more sensitive NAAT (NS) substantially decreased alternative-specimen yield of positive samples. NP swabs remain the gold standard for diagnosis of SARS-CoV-2, although alternative specimens are promising. Much remains unknown about the impact of variations in specimen collection, processing protocols, and population (pediatric versus adult, late versus early in disease course), such that head-to-head studies of sampling strategies are urgently needed.

KEYWORDS
COVID-19, molecular diagnostic, nasal swab, nasopharyngeal swab, oropharyngeal swab, saliva

Testing for SARS-CoV-2 (severe acute respiratory syndrome coronavirus 2, the etiologic agent of COVID-19) has preferentially utilized nasopharyngeal (NP) sampling with flocked swabs. This sampling method is presumed to have the highest diagnostic yield, as evidenced by its use as a reference method by the U.S. Food and Drug Administration (FDA) (1). However, NP sampling requires professional collection (a major limitation given the strained health care system in almost all parts of the world) and protective equipment which, similar to the flocked swabs themselves, is in short supply. In addition, NP sampling is uncomfortable, limiting patients’ willingness to come forward for testing, especially if asymptomatic.
If large-scale testing of symptomatic and asymptomatic patients is to become a reality, innovation in sampling is as important as innovation in testing. Innovation in sampling requires consideration of sampling methods (e.g., choice of swab, choice of sample type) as well as self-sampling.

As of October 2020, interim guidance from the U.S. Centers for Disease Control and Prevention (CDC) (2) recommends upper respiratory tract testing with any of the following specimens: NP swab, NP wash/aspirate, nasal wash/aspirate, oropharyngeal (OP) swab, nasal midturbinate (MT) swab using a flocked tapered swab, an anterior nares (AN) nasal swab using a flocked or spun polyester swab, or a saliva specimen obtained by supervised self-collection. The FDA, in contrast, states that NP, OP, MT, and AN swab samples are appropriate for clinical testing, and that “more data are necessary to better understand the performance when using specific saliva collection devices or other specimen types for COVID-19 testing” (1).

Saliva is the subject of the largest body of research on alternative specimen types, as well as the specimen of choice for numerous companies putting high-volume testing programs in place. Certain regions of the world, such as Hong Kong, have already adopted saliva in their mass screening protocols (3). The pathophysiological rationale for saliva sampling is based on the angiotensin-converting enzyme II (ACE-2) being the cellular receptor for SARS-CoV-2 (4), similar to SARS-CoV (5). High ACE-2 receptor expression in oral mucosa and salivary glands has been recently demonstrated (6, 7). Salivary gland duct epithelial cells previously were identified as a target for SARS-CoV in a rhesus macaque model (8). One study detected SARS-CoV-2 in saliva collected via expression directly from the salivary gland duct (9). These findings suggest that saliva may be a suitable and high-yield diagnostic sample for SARS-CoV-2 testing based on local viral replication, in addition to the possible mixing in saliva of lower and upper respiratory tract fluids that can carry virus. Saliva offers several advantages as it is a non-invasive sample type that can be self-collected, thus decreasing infectious risk to medical personnel, use of personal protective equipment, and reliance on equipment subject to supply shortages such as nasal, OP, or NP swabs.

OP (throat) swabs have been in wide use for SARS-CoV-2 diagnosis since the beginning of the pandemic (3). Consensus is lacking on both the best collection approach and nomenclature for oral specimens. The terms “oropharyngeal” and “throat” have been used in the literature, and therefore we describe and group them here as “oropharyngeal” or OP swabs. No comprehensive evaluations of their performance compared to other methods exist. OP swabs are less specialized than NP swabs and thus OP samples can be collected with a broader range of swab products. While the FDA recommends that OP swabs be collected by a health care professional (1), some have suggested that self-swabbing might be possible.

Nasal swabs (NS), another important alternative specimen type, also have the advantages of increased comfort and possible self-collection. They have been classified into two types anatomically. (i) Nasal midturbinate (MT) swabs, also called deep nasal swabs, are defined by the CDC (2) as flocked/tapered swabs performed while tilting the patient’s head back 70 degrees and inserting the swab less than one inch (about 2 cm into the nostril) until resistance is met at the turbinates before rotating the swab several times against the nasal wall. (ii) Anterior nares (AN) swabs are defined without a head tilt and inserting the entire swab at least 0.5 in. (1 cm) inside the nostril (naris) and sampling the membrane by various methods, including rotating the swab in place or around the inside wall of the nostril multiple times, and/or leaving in place for 10 to 15 s. The current “lower nasal swab” protocol sanctioned by the FDA specifies swab insertion “until you feel a bit of resistance” and thus matches the MT depth defined by the CDC, though swab type is not specified (10). The CDC and FDA suggest swabbing both nares for sampling.

A large body of literature on the yield of these alternative sample types for nucleic acid amplification testing (NAAT) has been created, but a comparative systematic summary of the relative performance of these alternative specimen types, compared to NP swabs, is lacking.
swabs, is missing and is needed to inform decision makers. Furthermore, the data for
collection of these alternative specimens (1, 2) have not been systematically evalu-
ated. Here, we aim to clarify the performance of alternative specimen types for diag-
nosis of SARS-CoV-2 by systematically reviewing and meta-analyzing the literature on
this topic available through October 2020.

**METHODS**

This systematic review and meta-analysis is reported in accordance with PRISMA
guidelines (see the supplemental material for the PRISMA checklist). The protocol for
this work was registered in the International Prospective Register of Systematic
Reviews (PROSPERO) (identifier: CRD42020214660).

**Search strategy, information sources, and eligibility criteria.** We performed a
comprehensive search of the Pubmed/MEDLINE and Google Scholar databases, as well
as the preprint servers medRxiv and bioRxiv, to identify relevant studies from 1 January
2020 until 1 October 2020. Only English language articles were allowed. An example
search strategy is provided in the supplemental material. Additional studies were
retrieved by screening the reference lists of the included articles and from archives of
the reviewers. We excluded papers from the same hospital with overlapping inclusion
dates, to avoid including patients more than once and thus minimize bias in the data
(11). Cross-sectional, case-control, and cohort studies and randomized controlled trials
were included independent of the number of specimens tested. Conference proceed-
ings and abstracts were deemed ineligible. Participants of all age groups with pre-
sumed SARS-CoV-2 infections, in all settings, were included. We included only papers
utilizing NAAT for SARS-CoV-2 detection.

**Data extraction.** Two reviewers assessed all articles (R.A.L. and J.C.H.) independ-
ently and disagreements were resolved with the input of a third investigator (N.R.P. or
C.M.D.).

We compared alternative sampling to NP sampling. We extracted only data for posi-
tive NAAT results on at least one sample type and only when sampling methods being
compared were performed synchronously. If patients were tested serially over time and
data could not be extracted for specific time points, we excluded the study. Asymptomatic, symptomatic, and unspecified symptomatology patient cohorts were
included. If multiple different RT-PCR tests were performed on one sample within one
study, we utilized the data from the best-performing assay (highest positive detection
rate) for a sample type. If a study contributed data to more than one analysis (e.g., two
different alternative sample types in one study, each compared to NP swab), it was
considered two or more data sets. If a combined NP/OP sample was used, and no data
were available for NP sampling alone, we included the study, using NP/OP as the com-
parator. For each specimen type, in addition to data on test performance, we extracted
data for factors likely to affect test performance, as detailed below. Data on throat or
gargle washes were not included in this meta-analysis (12, 13). We obtained the limit
development (LOD) from studies by direct report within the study when available, and
otherwise by manufacturer claims (package insert if available; Table S1 in the supple-
mental material lists how LOD was ascertained by study).

Our study retrieval process is depicted in Fig. S1. Records were organized using a
reference manager (Zotero Version 5.0.89, George Mason University).

**Assessment of methodological quality.** The Quality Assessment of Diagnostic
Accuracy Studies-2 (QUADAS-2) tool, a validated quality assessment tool for diagnostic
studies, was used to assess the included studies’ risk of bias (14). The four domains
assessed for risk of bias and applicability include: (i) participant selection; (ii) index test;
(iii) reference test; and (iv) flow and timing.

**Data analysis.** Under the assumption that (i) we do not know in advance which
specimen type performs best for SARS-CoV-2 detection and (ii) false positives are very
infrequent for NAATs performed in qualified laboratories, we focused only on individu-
als positive for at least one sample type and report on agreement between samples
rather than measures of sensitivity and specificity. Specifically, for each study, we
included all individuals with paired specimens who had at least one positive specimen as the denominator, and calculated (i) % positive Alt (percentage of individuals positive by the alternative specimen type [e.g., saliva, NS, or OP swab]); (ii) % positive NP (percentage of individuals positive by the NP specimen [or NP/OP in a minority of cases]); and (iii) % dual positive (percentage of individuals positive for both the alternative specimen and NP specimen).

We present results of the systematic review in forest plots. Meta-analyses were only performed when there were at least 4 primary studies and at least 20 patients per study. The % positive Alt, % positive NP, and % dual positive for pooled, overall, and in subgroups were estimated using a logistic-normal random-effects model (via the ‘metaprop_one’ command with Stata [15]). We applied the Freeman-Tukey double arc-sine transformation to stabilize variances and score 95% intervals were computed. Heterogeneity was measured by the inconsistency index (I²), which describes the percentage of variation across studies due to heterogeneity rather than chance. To investigate possible contributors to heterogeneity, we present subgroup analyses by (i) site of sampling; (ii) swab material (flocked, unflocked); (iii) sampling procedure; (iv) professional versus self-sampling; (v) specimen processing/NAAT (including dilution, nucleic acid extraction procedure, and assay used); (vi) populations (e.g., pediatric, asymptomatic); and (vii) symptom duration prior to testing. Due to heterogeneity in subgroups and small sample size, we report differences between subgroups descriptively but did not perform direct statistical testing between subgroups. The robustness of the meta-analysis to publication bias was assessed by the symmetry of funnel plots.

All analyses and graphs were performed using Stata 15.1 (StataCorp, Texas) and GraphPad 8.5 (Prism, SanDiego).

RESULTS

Our search yielded 1,253 unique citations, of which 25 were included in the analyses for saliva, 11 for NS, 6 for OP, and 4 for OP/NS swabs. Reasons for exclusions of studies are laid out in Fig. S1.

The studies we included in the meta-analysis overall had a moderate to high risk of bias according to QUADAS-2 (Table S2, Table S3), with studies on NS having a low to moderate risk (Table S4). Many studies did not specify patient selection methodology (random or consecutive). Many studies were also composed of cohorts of known positives (case-control) that were then retested with paired specimens and therefore at a high risk of a selection bias, limiting applicability of results to the general screening population.

Saliva. In total, 25 studies (16–40) met our inclusion criteria to assess saliva as an alternative sample type for SARS-CoV-2 diagnosis. The studies cumulatively included 4,528 paired saliva and NP swab specimens, although two studies used a combined NP/OP swab as a comparator (24, 39). An additional 13 studies (3, 8, 41–51) described a performance estimate for saliva as a specimen type, but these studies were excluded from the meta-analysis as the saliva and NP samples were either not collected synchronously, or multiple paired specimens were taken from the same set of patients and the data could not be extracted for a unique patient/time point.

Across the 25 studies, we found that the % positive saliva (88%, 95% confidence interval [CI] 81 to 93%) was lower than the % positive NP (or NP/OP), although not substantially different (94% [95% CI 90 to 98%]). The % dual positive was noticeably lower than either specimen type alone (79%, [95% CI 71 to 86%]) (Fig. 1), indicating relatively poor agreement. Considerable heterogeneity was also detected (I² 88.6%). This heterogeneity was likely attributable to the variation in study procedures and patient population between studies, as outlined in Table S5. Notably, there were no head-to-head studies for any of the comparisons described below for differences in collection, processing, and populations.

Saliva collection protocols for included studies were assessed for differences with respect to (i) asking patients to cough or clear their throat before submission
of sample (likely mixed sputum and saliva specimen or deep throat saliva specimen) or (ii) requesting the patients submit “drool” or “spit.” While some authors (42) have hypothesized that capture of posterior oropharyngeal saliva or mixed sputum/lower respiratory specimen is important for diagnostic sensitivity, we did not find a considerable difference in performance, although % positive saliva was

![Summary forest plot of individual studies assessing saliva.](image-url)
higher for studies (33, 34, 37, 40) that specified cough or deep throat saliva specimen versus studies that did not specifically ask for this (94% [95% CI 87 to 99%] versus 86% [95% CI 78 to 92%]) (Fig. 2). For NP samples in these two groups, the % positive detection was similar (89% [95% CI 60 to 100%] versus 95% [95% CI 93 to 97%]). Notably, many studies that supported the hypothesis of a coughing or deep throat saliva being better than drool/spit sample were excluded from the meta-analysis due to nonsynchronous sample collection or repeat testing on the same set of patients (3, 41–43).
Other differences in sampling between studies included avoidance of eating, drinking, or brushing teeth before specimen submission (typically 30 min to 2 h). Protocols that specified avoidance of eating, drinking, or brushing teeth prior to saliva collection (22, 23, 26, 31–34, 40) had higher % positive saliva, although the difference was not substantial (91% [95% CI 86 to 95%] versus 86% [95% CI 79 to 92%]) (Fig. 2). For NP sampling in the same two groups of studies, the difference was minimal (94% [95% CI 80 to 100%] versus 95% [95% CI 92 to 97%]). A few studies specifically requested morning submission (33, 40) in addition to avoidance of food/drink and similar saliva performance was identified between protocols that specified morning submission versus those that did not (92% positive saliva [95% CI 88 to 96%] versus 87% [95% CI 79 to 93%], respectively). However, for NP sampling in the two groups of protocols, the % positive NP was lower with morning submission (66% positive [95% CI 60 to 72%] versus 96% positive [95% CI 94 to 97%]), but this was largely driven by one study (33) in which NP swabs performed poorly (this was an outpatient cohort of patients who were asymptomatic at the time of collection and had received their diagnosis at least 1 week prior).

While many studies specified self-collected saliva (16, 17, 20, 25–27, 29, 30, 33, 34, 37, 39, 40), some studies described supervised collection (22, 28, 31, 35, 38). The % positive saliva was higher for self-collection than supervised collection, although the difference was not substantial (92% [95% CI 86 to 96%] versus 83% [95% CI 60 to 98%], respectively) (Fig. 2). Notably, only two studies (26, 33) reported using RNase P as a control for human material sampling adequacy.

There were also substantial differences in specimen processing, including variable dilution of the saliva specimen prior to NAAT, and use of protocols that directly input saliva samples into the NAAT without nucleic acid extraction. We found that even in the absence of dilution for saliva, the % positive could not match that of NP swabs. Studies utilizing undiluted saliva specimens (22, 26) had similar % positive saliva to studies utilizing diluted saliva (16, 17, 23–25, 27–32, 35, 37, 40) (Fig. 2) (92% [95% CI 86 to 97%] versus 89% [95% CI 81 to 95%]). All NP swabs were eluted in viral transport medium, and performance was similar in these two groups of studies (97% [95% CI 91 to 100%] for undiluted saliva versus 94% [95% CI 92 to 96%] for diluted saliva). In studies using diluted saliva, there was wide variation in dilution methods, with many groups not specifying the degree of dilution.

Studies that did not use a nucleic acid extraction step (19, 28) but instead directly input the saliva specimen into the amplification assay without any preprocessing showed substantially lower % positive saliva than studies that had an extraction step (60% [95% CI 49 to 70%] versus 89% [95% CI 83 to 94%]) (Fig. 2). In contrast, % positive NP swabs was not different between the two groups of studies (89% [95% CI 81 to 95%] versus 95% [95% CI 90 to 98%]), with all except one study (28) using a nucleic acid extraction for the NP swab sample (28).

While the majority of studies utilized reverse transcription PCR (RT-PCR) assays, one study (19) used reverse transcription loop-mediated isothermal amplification (RT-LAMP), and two studies used transcription-mediated amplification (TMA) (35, 36). The NAATs also differed over 10-fold in their limits of detection between studies, which has been previously described to affect diagnostic positivity (52). We found slightly higher, but not substantially different, percentages of positive detection in studies with a more sensitive/lower (<1,000 copies/milliliter [cp/ml]) limit of detection (LOD) (20, 21, 24, 34, 39) compared to studies with LOD of ≥1,000 cp/ml (23, 26, 27, 31–33, 51) (90% [95% CI 79 to 97%] versus 87% [95% CI 77 to 95%]). The differences were similar for NP swabs (100% [95% CI 97 to 100%] versus 91% [95% CI 71 to 100%]).

We also assessed if patient symptomatology could explain variable diagnostic performance between saliva and NP sampling. We found that only a few studies provided a direct comparison between asymptomatic (16, 19, 33) and symptomatic patient data (17, 18, 21–23, 25, 27, 28, 34, 35, 37, 39, 40) that could be parsed and extracted for analysis. We found that % positive saliva was similar between asymptomatic and
symptomatic patients (87% [95% CI 70 to 98%] versus 88% [95% CI 79 to 95%], respectively). The difference between % positive NP in asymptomatic versus symptomatic patients was much larger (symptomatic 96% [95% CI 93 to 99%] versus asymptomatic 73% [95% CI 47 to 93%]) (Fig. 2). These findings, however, were driven by one study (33) with superior saliva performance in asymptomatic patients (% positive saliva 93% [95% CI 88 to 97%] versus % positive NP 52% [95% CI 44 to 60%]). As discussed above, this study (33) included an outpatient cohort who had received their diagnosis at least 1 week prior. Notably, although the patient population is described as “asymptomatic,” it is unclear if this was just at the time of collection and they had symptoms closer to their initial diagnosis.

Another important question that we were not able to adequately address from the literature is the performance of saliva in pediatric populations. There are two studies (51, 53) that evaluated the performance of saliva in children, both of which showed worse performance compared to NP swabs (8/11 children positive by NP swab were positive by saliva in one study (51), and in the other study 53% of the children detected by NP swab were also positive by saliva ([53]). However, the timing of saliva collection versus NP swab collection in both studies was unclear, results for asymptomatic versus symptomatic children were not clearly distinguished, and sample processing methods were not clearly described.

Invalid test results were also not consistently reported across studies. Viscosity of saliva was highlighted in some studies as increasing errors in automated pipetting steps, necessitating dilution or biochemical pretreatment of samples (26, 35, 44).

Given that the fluctuation of viral load over time may differ between saliva and NP samples, and the time points at which patients present themselves for diagnosis may vary, we also assessed differences in diagnostic performance at different time points throughout the illness. Subgroup analysis of 6 studies with extractable data (25, 27, 28, 36, 39, 51) found that % positive saliva was overall lower at >7 days after symptom onset (74% [95%CI 62 to 85%]) compared to ≤7 days (89% [95% CI 73 to 99%]), which was also observed for NP swabs (91% [95% CI 82 to 98%] versus 99% [95% CI 90 to 100%]), respectively).

**OP swab.** We identified six studies that assessed OP versus NP swabs (54–59) and were suitable for meta-analysis. Given the paucity of data, subgroup analyses to assess differences in collection procedure, sample processing, and patient symptomatology were limited (Fig. 52). We found that % positive OP swabs was similar to % positive NP swabs (84% [95% CI 57 to 100%] versus 88% [95% CI 73 to 98%]) although % dual positive was only 68% (95% CI 36 to 93%), suggesting limited agreement (Fig. 3). Notably, the % positive NP estimate was unusually low in this group of studies, which is largely driven by one study (56) with a large gap in performance between % positive OP and % positive NP (86% [95% CI 65 to 97%] versus 41% [95% CI 21 to 64%], respectively) that was not observed in other studies. This study was unique in that paired samples were taken near the end of a hospitalization (unclear duration of symptoms) in a cohort comprised of known positives from prior NP swab RT-PCR-positive patients to help determine discharge eligibility.

Another outlier study in this data set was a cohort of symptomatic patients (55) in a Wuhan hospital early on in the pandemic with unusually poor performance of OP swabs (36% [95% CI 25 to 47%]) in comparison to NP swabs (88% [95% CI 79 to 94%]). There were not sufficient data in the OP swab meta-analysis to assess the impact of symptomatology or duration of symptoms on percent positives.

Only 3 studies specified flocking of swabs (54, 57, 59), and in all 3 cases unflocked oropharyngeal swabs were compared to flocked NP swabs. The percentage positive detection rate for these 3 studies was similar between OP and NP sampling (96% [95% CI 88 to 100%] versus 97% [95% CI 90 to 100%]). There were two studies that used health care worker-collected oral swabs (54, 57) and the positive detection rate for OP versus NP sampling was similar (97% [95% CI 89 to 100%] versus 96% [95% CI 87 to 100%], respectively) (Fig. 3). One study used
unobserved self-collected OP swabs and reported similar performance between sample types but in a limited sample set of only 12 positive patients (92% [95% CI 62 to 100%] for OP versus 100% [95% CI 74 to 100%] for NP). The 3 other studies (55, 56, 58) in this meta-analysis did not specify self-collection versus health care worker collection. There was one study (60) that compared self-collected versus lab technician-collected OP swabs and found that only 14/24 total positives were detected by self-collection versus 22/24 total positives for lab technician collected. This study was excluded from this meta-analysis, however, as the pairing of OP swabs to NP swabs was unclear.

Three studies that assessed oral swab specimens were excluded from this meta-analysis as they were not oropharyngeal. There was one pediatric study of buccal swabs in Singapore (61) where children underwent daily NP and buccal swabs; 9/11 children with SARS-CoV-2 detected by NP swab also at some point had positive buccal swabs. One study that described sampling of the anterior 2/3 of the dorsum of the tongue (62) found similar positive detection rates from “tongue” swabs (46/51 total positives) and NP swabs (49/51 total positives). Another study asked patients to cough prior to sampling oral fluid in cheeks, gums, hard palate, and tongue (63), and reported that detection was slightly higher (26/29 total positives) than NP swabs (23/29 total positives).

**AN and MT swabs.** We identified 11 studies using either AN or MT swabs that could be combined for pooled meta-analysis (32, 35, 54, 59, 62–68) with NP swabs as the reference sample type, although one study (59) used a combined NP/OP swab for reference. We found that the % positive NS (either AN or MT) (82% [95% CI 73– 90%]), was substantially lower than % positive NP swab (98% [95% CI 96 to 100%]) as well as % dual positive (79% [95% CI 69 to 88%]), suggestive of limited agreement between the two sample types (Fig. 4). Considerable heterogeneity was again detected between studies ($I^2 = 87\%$), which we attributed to differences in study procedures. Accordingly, we assessed performance by collection protocol, self-collection/supervision/health care worker collection, sample processing, and patient symptoms (Table S6). Notably, there were no head-to-head studies for any of the comparisons.

![Summary forest plot of individual studies assessing oropharyngeal swabs.](image)
The pooled estimates of % positive NS via AN (35, 62) versus MT swabs (using the depth of insertion to classify AN versus MT sampling, per CDC description) were similar (54, 59, 63, 66–68) (90% [95% CI 84 to 94%] versus 84% [95% CI 65 to 97%]) (Fig. 5). The % positive NP was similar between groups (99% [95% CI 95 to 100%] versus 97% [95% CI 92 to 100%], respectively). There were only two studies that assessed AN swabs and, notably, one of the studies (62) also compared AN to MT sampling (AN performed before MT). Performance was similar between AN and MT swabs (48 out of 51 total positives for AN versus 50 out of 52 total positives for MT). In this study, both the AN and MT swabs were collected from both nares, and the AN swab was unflocked, whereas the MT swab was flocked.

We found that a more sensitive assay (LOD < 1,000 copies/ml) (64, 65, 68) resulted in worse performance of NS in comparison to assays with LOD ≥ 1,000 copies/ml (32, 54, 59, 63, 66, 67) (61% [95% CI 40 to 79%] versus 85% [95% CI 82 to 91%]), while % positive NP were similar (99% [95% CI 93 to 100%] versus 97% [95% CI 92 to 100%], respectively). This may reflect lower viral burden in the midturbinate/anterior nares region than the nasopharynx, resulting in lower performance in comparison to NP swabs that is only evident when using a highly sensitive (LOD < 1,000 copies/ml) assay. Notably, the discordant paired samples described in the studies also were found to have lower viral loads than concordant pairs.

FIG 4 Summary forest plot of individual studies assessing nasal swabs.
Two of the studies (64, 68) with the worst NS performance used a more sensitive assay and also compared unflocked NS to flocked NP swabs (positive NS detection 48 to 56%). There were 3 other studies (35, 54, 59), however, that also used unflocked NS compared to flocked NP swabs and reported higher detection (83 to 100%). Ultimately, NS studies that specified that both NS and NP swabs be flocked (63, 66) had a substantially higher % positive NS than studies where an unflocked NS was compared to a flocked NP swab (35, 54, 59, 64, 68), although confidence intervals were wide and overlapping (90% [95% CI 81 to 97%] versus 77% [95% CI 55 to 93%]) (Fig. 5). This finding

**FIG 5** Summary forest plot of subgroup data from nasal swab sampling for different clinical populations, and collection as well as processing procedures.
may have been partially driven by the two studies using the lower LOD assay. The % positive NP between these two groups was similar (97% [95% CI 90 to 100%] versus 97% [95% CI 94 to 99%]). One study (62) collected unflocked AN swabs, flocked MT swabs, and unflocked NP swabs for comparison and reported similar performance of all three specimens, as described above. However, use of a nonflocked swab for the NP sampling may have decreased its sensitivity and artificially increased the sensitivity of MT and AN sampling.

Only 4 studies (35, 59, 62, 68) specified that NS were performed before NP swabs and although % positive NS was higher in comparison to studies that did not specify the swab order, there was not a substantial difference (85% [95% CI 58 to 100%] versus 81% [95% CI 74 to 88%]) (Fig. 5). The % positive NP was the same regardless of NS order (98% [95% CI 95 to 100%]). Surprisingly, NS specimens collected from both nares (35, 54, 62, 64, 67, 68) seemed to perform worse in comparison to swabs collected from a single nostril (59, 63, 66), although the difference was not substantial (77% [95% CI 59 to 91%] versus 93% [59 to 91%]) (Fig. 5). The % positive NP was again similar in these two scenarios (95% [95% CI 95 to 99%] for both nares versus 97% [95% CI 83 to 100%] for one nare). Notably, this finding may again have been driven by the two studies (64, 68) utilizing a more sensitive (<1,000 cp/ml) NAAT assay (both with poor detection by NS and performed with swabs collected in both nares), a factor which we described previously as associated with a lower NS detection rate.

We also found that unsupervised self-collected NS specimens (35, 59, 62) had higher percent positives in comparison to swabs collected by health care workers (54, 64, 67, 68) (93% [95% CI 85 to 98%] versus 68% [95% CI 47 to 86%]) (Fig. 5). In only one study (63) was the patient’s self-collection supervised (92% [95% CI 74 to 99%]). Professional NP sampling performance was similar between the NS self-collection and HCW-collection groups (99% [95% CI 96 to 100%] versus 96% [95% CI 92 to 99%]). Notably, the same two studies (64, 68) showing the worst NS performance (and using more sensitive assays) were in the health care worker-collected group. Additionally, only two studies (62, 63) reported use of RNase P as a human sampling control.

We could not find any studies that specifically assessed only asymptomatic patients with NS, although multiple studies used mixed populations and a few studies specified that all of the cohort was symptomatic. Studies of only-symptomatic patients (35, 62, 64–67) had a similar % positive NS rate to studies of mixed or unspecified patients (82% [95% CI 73 to 91%] versus 83% [95% CI 62 to 97%]) (Fig. 5). The % positive NP was similar between groups (99% [95% CI 98 to 100%] versus 96% [95% CI 88 to 100%]). There were two studies (63, 67) with extractable data to assess the impact of symptom duration prior to testing, and a lower yield after 7 days was found although the difference was not substantial (76% [95% CI 60 to 87%] for >7 days symptoms versus 88% [95% CI 74 to 95%] for ≤7 days) (Fig. 5). The % positive NP was similar in both groups (100% [95% CI 86 to 100%] versus 100% [98 to 100%]).

Again, data were limited on pediatric populations when using nasal swabs. There are two pediatric studies of nasal samples; one study (69) described NS to outperform OP swabs in 56 paired samples from 11 pediatric patients, with cycle threshold (CT) values lower in NS versus OP for all 11 first paired samples. This study had to be excluded due to repeat sampling, as we were not able to extract unique patient data from different time points. The other study (70) was not a nasal swab study, but described NP aspirates to outperform NP swabs for detection (% positive for NP aspirates was 88% in comparison to 51% for NP swabs), though details of methods provided were minimal and repeat sampling on patients occurred.

Combined OP/NS as a specimen type. Four studies (59, 71–73) evaluated combined oropharyngeal and nasal swabs in comparison to NP swabs (Fig. 6). Three of these studies (71–73) specified that a single swab was used for collection of an OP/NS sample, whereas in one study it was unclear if two separate swabs were used for OP and NS sampling and the results compiled (59). Two of the studies (59, 73) used MT swab depth, while one used AN sampling (72), and one was unspecified (71). Three of
the studies were health care-collected swabs, in which the OP sampling was performed prior to NS (71–73), and only two of these studies specified swabbing both nares (72, 73). Two studies used flocked swabs (71, 73), while the others used unflocked swabs. The LOD of the assay in copies/ml was only available in one study (59) and was >1,000 cp/ml. Two of the studies (59, 71) found that the percent positive detection of the combined swab specimen was greater than percent positive detection for the reference NP swabs. Pooled detection estimates were similarly high between the combined swabs and NP swabs with the same % positive estimate (97% [95% CI 90 to 100%]), although agreement between the two methods was less (90% dual positive [95% CI 84 to 96%]).

Assessment of publication bias. Visual inspection of a funnel plot of the study data versus standard error shows substantial asymmetry and therefore suggests publication bias for all alternative sample types skewed toward publication of positive findings (i.e., strong performance of alternative specimens) (Fig. S3).

DISCUSSION

This systematic review and meta-analysis synthesizes a large number of studies comparing alternative sample types to NP swabs for SARS-CoV-2 detection by NAAT. While all 3 sample types independently seemed to capture lower % positives (nasal swabs 82% [95% CI 73 to 90%], OP swabs 84% [95% CI 57 to 100%] and saliva 88% [95% CI 81 to 93%]) in comparison to NP swabs, combined OP/nasal swabs in 4 studies, interestingly, had the same % positive detection rate as NP swabs (97% [95% CI 90 to 100%]) (Fig. 6).

For saliva specimens, we found slightly lower performance compared to NP swabs overall (88% [95% CI 81 to 93%] versus 94% [95% CI 90 to 98%]), and only the absence of nucleic acid extraction resulted in a substantially lower rate of detection. One study (44) had to be excluded from the meta-analysis due to nonsynchronous collection of NP and saliva samples. This study is of interest nevertheless, as it tested different RT-PCR platforms on the same set of saliva specimens using 3 different extraction-free commercial RT-PCR assays against a standard RT-PCR assay with extraction and reported that 79, 81, and 52 positive specimens, respectively, were detected out of a

FIG 6 Summary forest plot of individual studies assessing oropharyngeal/nasal swab sampling.
total of 84 positive specimens detected on the standard assay, indicating that choice of extraction-free assay matters.

Self-collection, coughing or deep throat saliva, and avoiding food, drink, or tooth brushing resulted in >5% increased positive saliva detection rates, although the difference was not substantial. Collection of saliva greater than 7 days after symptom onset also resulted in >10% lower detection, although the difference was not substantial. Viscosity has been described qualitatively in multiple studies as a challenge in utilization of saliva as a specimen type, but invalid rates were not documented in many studies; we note that invalid rates are a critical parameter and should be consistently reported.

We found that OP swabs seemed to perform similarly to saliva and NP swabs, but these estimates were highly affected by one study (56) where samples were collected near the end of a hospitalization for discharge purposes and the % positive NP was unusually low in comparison to the rest of the literature. Overall, the data argue for provider-facilitated collection and against self-collection of this sample type.

For NS, the literature to date supports that these perform worse than NP swabs, although these findings were largely driven by two studies (64, 68). There was no substantial difference between AN and MT swab detection, and we note that AN/MT swab collection protocols may have overlapped in practice; we defined AN versus MT based on depth of insertion, as the swab type was not always specified. We hypothesize that the difference in NS to NP performance was driven by use of a particularly sensitive (<1,000 cp/ml LOD) assay for two of the studies. We observed that the discordant samples (NP+/NS−) typically had low viral loads on the NP sample. While this may reflect differences in viral burden anatomically between NP and AN/MT sampling, the clinical significance of this difference remains to be determined.

We hypothesize that nasal swab performance is likely to be highly dependent on collection procedure, which has in turn evolved over time and a substantial amount of data remains unpublished (Nira Pollock personal communication). Self-collection and single-nare sampling trended toward higher, although not substantially different, % positive detection, although these results were largely driven by the two studies with poor NS performance (64, 68). Detection was also lower after >7 days of symptoms, although this difference was not substantial. There remain unresolved questions on the best-performing swab material (spun polyester, foam, or rayon) and sample transport (buffer versus dry swab) that could not be addressed in this study due to limited data (many studies did not report the swab product or elution details). We also still do not fully understand the impact of flocking on this specimen type. While studies using flocked nasal swabs had a slightly higher % positive detection in comparison to unflocked swab studies, the unflocked swab group included two studies using a lower LOD assay, and the 3 studies that used a less sensitive assay had near 90% detection (similar to the flocked group).

OP/NS samples had, surprisingly high performance compared to NP swabs; while this sample type likely requires an operator for collection, additional studies are warranted to understand acceptability in adult and pediatric patients.

Overall, much remains unknown about the impact on diagnostic sensitivity of variations in specimen collection and processing protocols, and performance in specific key subpopulations (asymptomatic versus symptomatic, pediatric versus adult, late versus early testing from symptom onset). While we assessed aggregate data from different studies to gain insight into these variables, a limitation of this meta-analysis is that true comparison is precluded in the absence of head-to-head studies. Furthermore, while there are trends we observe in our subgroup analyses, these findings may be population related and should be interpreted with caution. Timing of sampling from symptom onset was also quite variable (collection occurred within days to weeks in some studies), and was inconsistently reported, which likely had a major impact on diagnostic performance, given decreasing viral load over time. Head-to-head studies are urgently needed of flocked versus unflocked swabs (and specialized versus unspecialized
swabs for MT collection), collected at different times in disease and with different sampling methods, and also in important subpopulations (e.g., children), to resolve the persistent uncertainty.

We note that the reporting quality of studies was low, STARD guidelines (74) were not consistently followed, and study bias was considered moderate to high on QUADAS 2. Lastly, in this study we chose to report the % positive alternative-specimen, % positive comparator-specimen, and % dual positives instead of the positive percent agreement (PPA). This decision was motivated by our presumption regarding the low rate of false positives using NAAT, and the potential for an alternative to yield more positive results than the comparator NP, which would otherwise not be considered.

In summary, while alternative specimens (particularly saliva and OP/NS samples) show promise, we find that the literature to date suggests that NP swabs are indeed the gold standard in comparison to alternative specimen types (saliva, OP swab, NS). We identify self-collected nasal swabs and saliva to have similar performance to health care worker-obtained specimens, which is also supported by a head-to-head comparison of self-collected AN versus professionally collected NP swabs utilizing an antigen rapid test for detection (75). We reiterate that the LOD of any assay will impact detection and centers should be aware of the increased possibility of false negatives with any sample type when using a less sensitive assay. Given the promising results of combined oropharyngeal and nasal swab studies, more studies on alternative specimen combinations would be useful. Lastly, we encourage future studies to provide more clarity about exact details of collection procedures, specific swab shape and materials used, sample processing methods (dilution, extraction, storage, transport), and the NAAT assay utilized (including LOD), allowing the field to clearly define the trade-offs required to sufficiently bring SARS-CoV-2 testing to scale.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

SUPPLEMENTAL FILE 1, PDF file, 0.6 MB.

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REFERENCES


