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The performance of wearable sensors in the detection of SARS-CoV-2 infection: a systematic review

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1 Introduction

On Dec 31, 2019, WHO recognised the emergence of SARS-CoV-2, a novel virus in the coronavirus family.1 Since then, the outbreak of illness caused by the SARS-CoV-2 virus (COVID-19) has become a global pandemic, causing more than 458 million cases and 6 million deaths, until March, 2022.2

A key strategy for containing the COVID-19 pandemic has been the rapid identification and contact tracing of infected individuals.3–5 RT-PCR constitutes the gold standard for diagnostic testing of COVID-19.6–7 Despite developments in rapid testing, the timing of testing in relation to stage of infection hinders public health efforts to control the virus.8 On average, from SARS-CoV-2 infection to symptom onset takes 6 days, although the incubation period can be as long as 18 days.9 The viral load from the upper respiratory tract increases during the incubation period, reaches a peak around symptom onset, and then gradually declines.10 Many national health guidelines recommend testing for the general population after symptom onset, or a few days after suspected exposure to the virus, regardless of symptoms, to limit false-negative test results.11–14 However, viral load could be sufficiently high enough for transmission before people have symptoms or qualify for testing.15,16

COVID-19 remains difficult to distinguish from other respiratory illnesses on the basis of reported symptoms alone. Many common COVID-19 symptoms (eg, fever and cough) overlap with other influenza-like illnesses.17 Some patients with confirmed COVID-19 report symptoms uniquely associated with the virus (eg, anosmia), but such symptoms rarely appear early in the disease.18 Furthermore, 20–30% of individuals infected with SARS-CoV-2 never develop symptoms.19–21 The US Centers for Disease Control and Prevention report that presymptomatic or asymptomatic people account for half of SARS-CoV-2 virus transmissions.22–24 To reduce transmission rates in the general population, identifying SARS-CoV-2 infections before or in the absence of symptom onset is crucial. A range of non-invasive, commercially available physiological monitors (ie, wearable devices) could help in detecting presymptomatic and asymptomatic infections and controlling the pandemic. Because of rapid technological advancements, relatively subtle fluctuations in physiological parameters such as body temperature, respiratory rate, heart rate, heart rate variability, skin perfusion, and oxygen saturation (SpO2) can be measured by sensors commonly found in smartwatches, smart rings, and fitness trackers. Fever remains one of the most commonly reported COVID-19 infection symptoms;25 thus, the inclusion of thermometers sensors on an increasing number of wearable devices, despite their reliance on sensors worn on distal body parts, might render them suitable to detecting SARS-CoV-2 infection. Of note, peripheral temperatures measured by wearable devices have shown greater sensitivity than oral measurements in detecting subtle temperature shifts (eg, ±0.2°C).26 With regard to the COVID-19 pandemic, wrist temperatures have been found to be equally stable and less susceptible to environmental influences than forehead temperatures.26 Calls for additional research on the role wearable devices...
could serve in the early and comprehensive detection of SARS-CoV-2 infections have emphasised their potential ability to inform population and individual health responses to the pandemic. Several studies, mostly of retrospective design, have shown the feasibility of wearable devices in indicating the presence of SARS-CoV-2 infection by monitoring one or more physiological parameters, but an overview of the evidence is not yet available.

In this systematic review, we aimed to summarise and assess the added value of wearable devices in the detection of SARS-CoV-2 infection within the adult population (ie, those 18 years and older). Our primary question regards the current state of evidence on the diagnostic accuracy of statistical and algorithmic models using wearable sensor data. We also consider the time from detection to symptom onset and which physiological parameters provide the best indication of a subclinical or symptomatic SARS-CoV-2 infection.

Methods

Search strategy and selection criteria
We conducted our systematic review in line with our protocol and report our findings according to PRISMA recommendations. We initially searched the literature between Dec 17 and Dec 21, 2020, on the electronic databases PubMed (MEDLINE), Embase, Web of Science, Cochrane Central Register of Controlled Trials (known as CENTRAL), International Clinical Trials Registry Platform, and ClinicalTrials.gov. As the use of wearables to identify SARS-CoV-2 infections remains an ongoing area of research, we also searched preprint repositories (medRxiv and bioRxiv) for non-peer-reviewed studies between Dec 17 and Dec 21, 2020. We manually searched the reference lists of articles and reviews included for inclusion criteria underwent data extraction to obtain study-level information on participant demographics, study design and setting, sample size, the type of wearable device and its sensors, reference test, definition of key model parameters and features, and performance metrics (eg, area under the curve [AUC] and other test statistics). We contacted all corresponding authors to discuss missing data and areas of uncertainty. Finally, we assessed the risk of bias for each study’s primary outcomes using an adapted version of the National Institutes of Health Quality Assessment Tool for Observational Cohort and Cross-Sectional Studies. Per our protocol, a meta-analysis of the results could not be done, given the heterogeneity in approaches and outcomes.

Results

The first database search, done on Dec 17–21, 2020, identified 1601 records with an additional four articles retrieved from manually screening review reference lists. The second search, conducted on March 8–9, 2021, found an additional 574 records, and the third search, done on July 27, 2021, found an additional 1691 records, resulting in 3196 unique records overall, after deduplication. After title and abstract screening, 173 articles were retained for full-text review, of which 12 fulfilled our inclusion and exclusion criteria (appendix p 7). All studies were observational, and seven were strictly retrospective, although some researchers implemented control procedures, no RCTs were reported. Our searches also identified 12 study protocols, including two RCTs.

Eight protocols were recorded in online registries; one was a preprint, and three were published (appendix pp 8–9). During extraction, we contacted the corresponding authors for studies with missing data and received replies from six of the 12 research teams. We compiled the key characteristics for the 12 studies included in this systematic review (table 1; see appendix pp 12–14 for a detailed description). Most studies recruited active users of wearable devices with a
| Study design/Population and study setting | Wearable device | COVID-19-positive sample size (n)/Total analysed sample size (n) | Race or ethnicity n (%) | Reference standard | Timing of the detected deviation compared to SD (days)* | Algorithm or statistical model | Key features in the best performing model | Training set, n (%) | Test, validation (internal or external), or comparison set, n (%) | Key findings for the best performing model |
|------------------------------------------|----------------|---------------------------------------------------------------|------------------------|-------------------|----------------------------------------------------|-------------------------------|----------------------------------------|------------------|-------------------------|---------------------------------------------|
| Bogu and Snyder (2021)                  | Fitbit         | 25/106                                                        | NR                     | Self-reported COVID-19 diagnosis confirmed with a physician note | SO -6·94 to +5·12 | Long-term short-term memory networks-based autoencoder for anomaly detection (known as LAAD) | Changes in resting HR from baseline | NR | NR | Positive predictive value 0·91 (95% CI 0·85–0·967); sensitivity 0·36 (95% CI 0·23–0·487); F-beta (0·1) 0·79 (95% CI 0·69–0·88); abnormal HR lasted longer in the COVID-19-positive cohort; more COVID-19-positive cases had >1 day of abnormal HR |
| Cleary et al (2021)                     | Fitbit Inspire HR and the Apple Charge 3 Watch | 22/105                                                          | NR                     | Self-reported SARS-CoV-2 test | NA | Binary classifier | Resting HR, sleep duration (min), and total step count | 105 (100%) | NA | Activity data: AUC 0·75 (95% CI 0·63–0·87); all sensor data: AUC 0·75 (0·62–0·89) |
| Hassantabar et al (2020)                 | Empatica E4, pulse oximeter, and blood pressure monitor | 57/87                                                          | NR                     | PCR test upon hospital arrival | ND | Deep neural networks | Galvanic skin response, SpO2, blood pressure, and questionnaire data (eg, on symptoms, presence of chronic lung diseases, and whether participants are immuno-compromised) | 52 (60%; 18 healthy, 16 asymptomatic, and 18 symptomatic) | 17 (20%) in test set (6 healthy, 5 asymptomatic, and 6 symptomatic); F1 (ie, the harmonic mean of precision and sensitivity) 98·2%; true positive 98·1%; false positive 0·8%; false negative 0% for symptomatic individuals; sensitivity 97·52%; specificity 99·16% |
| Hirten et al (2021)                      | Apple Watch Series 4 or 5 | 13/297                                                        | 73 Asian (24·6%), 29 Black (9·8%), 43 other (14·5%), 108 White (36·4%), 44 Hispanic ethnicity (14·8%) | Self-reported nasal PCR test | ND | Mixed-effect cosinor model | HRV (SD of normal to normal R–R intervals) including mean MESOR, acrophase, and amplitude | NA | NA | Shorter mean SD of normal to normal R–R intervals amplitude in participants positive for COVID-19 |
| Lonini et al (2021)                      | Unnamed throat-worn patch | 15/29                                                        | NR                     | Tested positive for COVID-19 | Logistic regression with elastic net regularisation | HRV, HR (SD of R–R intervals), respiratory rate, cough frequency, and walk cadence | 29 (100%); randomly sampled one walk sequence and one cough sequence with replacement five times per individual, repeated 100 times to estimate CI | Validation set was leave-one-subject-out cross validation | AUC ≥0·97 (95% CI 0·92–0·96); pre-walk HR higher in COVID-19-positive cohort; pre-walk respiratory rate higher in COVID-19-positive cohort; pre-walk HR lower for COVID-19-positive cohort; COVID-19-positive cohort walk slower |
| Study design / Population and study setting | Wearable device | COVID-19-positive sample/Total analysed sample size (n/N) | Race or ethnicity n (%) | Reference standard | Timing compared to SO (days)* | Algorithm or statistical model | Key features in the best performing model | Training set, n (%) | Test, validation (internal or external), or comparison set, n (%) | Key findings for the best performing model |
|-------------------------------------------|-----------------|--------------------------------------------------------|-------------------------|-------------------|-----------------------------|---------------------------------|----------------------------------------|-------------------|---------------------------------------------------------------|---------------------------------------------------------------|
| Miller et al (2020)                       | Observational and retrospective/Ambulatory, opt-in study of device users | WHOOP | 81/271 | NR | Self-reported SARS-CoV-2 test | ND | Gradient boosted classifier | $5$ respiratory rate-derived features | 57 (70%); COVID-19-positive with symptoms between March 14 and April 14, 2020 | 24 in validation set one (30%); COVID-19-positive with symptoms between April 14–June 6, 2020; 190 participants negative for COVID-19 in validation set two | Sensitivity 36·5%, specificity 95·3%, positive predictive value 73·8%, negative predictive value 80·6% for the test set; identified 20% of individuals positive for COVID-19 before SO, identified 80% of individuals positive for COVID-19 by SO +3 days |
| Mishra et al (2020)                       | Observational, prospective, and retrospective/Ambulatory, opt-in study of device users | Fitbit (Ionic, Charge 4, and Charge 3) | 32/120 | European (84·4%), 5 mixed or other (15·6%)§ | Self-reported COVID-19 (diagnosis confirmed with physician note) | SO –28 to SO +7¶ | Offline (HROS-AD, RHR-Diff) and online (CuSum) anomaly detection algorithms | The HROS-AD model included HR and step count as features; the RHR-Diff model included HR; the CuSum model included deviations in elevated residual resting HR | 32 (100% of participants positive for COVID-19) | 73 self-reported healthy participants in comparison set one; 15 participants not positive for COVID-19 in comparison set two | Median time to SO from elevated HR was 4 days, median HR increased by 7 beats per min following SO, step count decreased at onset of HR changes associated with COVID-19, sleep duration increased at onset of HR changes associated with COVID-19 when missing data was imputed, CuSum detected 63% of SARS-CoV-2 infections before SO in real-time |
| Natarajan et al (2020)                    | Observational and retrospective/Ambulatory, opt-in study of American and Canadian device users | Fitbit | 1257 | NR | Self-reported PCR test | SO –1 to SO +4¶ | Convolutional neural network | Body-mass index, age, sex, mean nocturnal respiratory rate, mean nocturnal HR during non-rapid eye movement sleep, HRV (RMSSD of nocturnal respiratory rate series), Shannon entropy of nocturnal respiratory rate series, and data from the day of examination and the 4 preceding days | 87% (random 70% split); 70:15:15 split performed five times, but cross-validation performed only once | 189 in test set (15%); 189 in cross-validation set (15%) | Sensitivity 25·9%, specificity 99·0%, AUC 0·77 (0·02), the 90% specificity model identified 40 (21%) of individuals positive for COVID-19 at SO –1, correctly identified 105 (56%) of individuals positive for COVID-19 at SO +4 |

(Continued from previous page)
| Study design / Population and study setting | Wearable device | COVID-19-positive sample/Total analysed sample size (n/N) | Race or ethnicity n (%) | Reference standard | Timing compared to SO (days)* | Algorithm or statistical model | Key features in the best performing model | Training set, n (%) | Test, validation (internal or external), or comparison set, n (%) | Key findings for the best performing model |
|-------------------------------------------|----------------|---------------------------------------------------------|-------------------------|-------------------|-----------------------------|-----------------------------|--------------------------------------|------------------|---------------------------------------------------------------|------------------------------------------------------|
| Nestor et al (2021)†† | Observational and retrospective (same data collection as used in Shapiro et al.†) / Ambulatory, opt-in study of device users | Fitbit | 204/32 198 | NR | Self-reported SARS-CoV-2 test or medically diagnosed influenza | Day of symptomatic infection (SO to symptom end) | Model 1 (wearable only data) was (1a) gradient boosted classifier and (1b) gated recurrent unit-decay; model 2 (survey only data) was gated recurrent unit; model 3 was paired gradient boosted classifier and gated recurrent unit | Model 1 included 48 features based on HR, steps, and sleep data; model 2 included survey data (daily symptom history and demographic covariates); model 3 included 48 features and survey data | 11 269 (35%); 35 7 55 50 split performed five times | 16 099 (50%) in test set one (prospective); 2415 (7.5%) in test set two (retrospective, held-out set); 2415 (7.5%) in validation set (retrospective) | Model 3 sensitivity 0.65 (95% CI 0.49–0.87); specificity 0.69 (95% CI 0.43–0.97); model 3 detects 63.5% of COVID-19-positive cases at SO (vs 47.7% for non-COVID-19 positive influenza like illnesses) |
| Quer et al (2021)†† | Observational and prospective / Ambulatory, opt-in study of American smart device users | Device-agnostic | 54/333 | NR | Self-reported COVID-19 test result | NA | Binary classifier | Resting HR, age, sex, cough, fatigue, decreased taste or smell, sleep duration (min), and total step count | 333 (100%) | NA | AUC 0.80 (95% CI 0.73–0.86); sensitivity 0.72 (95% CI 0.59–0.83); specificity 0.73 (95% CI 0.68–0.78); positive predictive value 0.35 (95% CI 0.29–0.41); negative predictive value 0.93 (95% CI 0.90–0.96) |
| Shapiro et al (2021)†† | Observational and retrospective digital cohort / Ambulatory, opt-in study of device users | Fitbit | 41/1352 | No American Indian or Alaskan Native, 4 Asian or Pacific Islander (9.8%), 3 Black or African American (7.3%), 4 Hispanic or Latino (9.8%), 3 preferred not to answer (7.3%), 4 unavailable (9.8%), 23 White (56.1%) | Self-reported COVID-19 diagnosis by a health-care practitioner | SO – 2 to SO + 2 | Multilevel model | Resting HR, week of flu season, day of the week, average activity level in participant’s physical state, and participant’s baseline activity level | NA | NA | Increased HR in COVID-19-positive cohort; increased sleep persisted for longer in COVID-19-positive cohort; COVID-19-positive cohort took fewer steps |

(Table 1 continues on next page)
self-reported retrospective SARS-CoV-2 infection; none of the studies tested participants for the presence of SARS-CoV-2 antibodies to detect mild or asymptomatic infections for which the participant had not sought diagnostic testing. Researchers commonly used historical information from long-term wearable use to examine changes in physiological parameters in the days before and after a patient’s diagnosis or symptom onset. The studies recruited predominantly from European and North American countries. Nine studies examined SARS-CoV-2 infection among the general public, whereas three enrolled health-care professionals.31,32,37 Three research teams characterised their studies as proof-of-concept studies.38–40 Four studies were pre-prints.31,32,36,40

The participant sample size (n=29 to 32198), sex ratio (17–70% male and 30–81% female), and mean ages (29–57 years) varied widely between studies. Information on ethnicity and race was collected and analysed in five studies,39,32,33,31,37,36 with only two studies recruiting a relatively diverse population.32,37

Various wearable devices were investigated across the 12 studies, with bracelet design constituting the most common style. Five studies examined physiological parameter changes exclusively32,31,33,35,40 or almost exclusively (99%) measured by Fitbit devices. Other, less commonly investigated wrist-worn devices included the WHOOP strap,30 the Apple watch,37 and the Empatica E436 (one study each). One study examined a smart ring, the Oura,39 whereas another study analysed data from an unnamed device worn on the user’s throat.38 The final study remained device-agnostic; most participants wore Fitbits (78–4%), but any device that paired with Apple HealthKit or Google Fit met eligibility criteria and was included.40 The 12 studies examined wearable device-measured physiological changes in respiratory rate,30,35,38,39 heart rate,35,36,33,35,38,40 heart rate variability,35,37,39 skin temperature,34,38 and movement31,33,31,34,38 (table 2; appendix pp 15–20).

Across studies, the research teams drew on diverse methods for examining wearable devices’ ability to detect SARS-CoV-2 infection (appendix pp 10–11). Nine studies used machine learning algorithms to identify how physiological data (supplemented by symptom reports in three studies31,33,34) could detect SARS-CoV-2 infection,31,33,34 including an anomaly detection auto-encoder,6 gradient-boosted classifiers,12,13 and deep,6 convolutional,13 or gated recurrent-unit14 neural networks. The remaining three studies used statistical analyses, such as mixed-effect models30,37 and Wilcoxon rank-sum tests.30 Although some studies examined differences between SARS-CoV-2 infection and other influenza-like illnesses,39,32,31,34 eight authors focused solely on SARS-CoV-2 infection. Nine studies built models to directly compare wearable data from patients positive for SARS-CoV-2 with healthy39,33,31,34 or SARS-CoV-2-negative controls.39,32,31,34

Eight studies considered intra-participant changes in baseline parameters as they progressed from uninfected to symptomatic to asymptomatic infection.32,31,33,31,37,36,40

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Table 1: Eligible studies using wearable devices to detect changes in physiological parameters among COVID-19 positive individuals

| Wearable device | Study design | Race or ethnicity n (%) | Number of participants | Reference standard | Timing compared to SO (days)* | Key features in the best performing model | Algorithm or statistical model | Timing compared to SO | Key findings for the best performing model |
|-----------------|-------------|-------------------------|------------------------|------------------|-------------------------------|----------------------------------------|-------------------------------|-------------------|-------------------------------------------|
| Outlining       | retrospective/observational | Asian (2%), White (78%), Hispanic or Latino (16%), Middle Eastern (2%), European (2%), Scandinavian (2%), Jewish (2%), South Asian (2%), unavailable (4%) | 120 | SARS-CoV-19 infection | 0 to 7 | Separate models for temperature, respiratory rate, HR, and HRV | SO to SO +7 Wilcoxon rank-sum test, Kruskal-Wallis non-parametric comparison | NA | Temperature increases around SO; respiratory rate increases after fever-based SO; HR increases after fever-based SO; HRV increases after fever-based SO |

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In general, algorithmic models for detecting SARS-CoV-2 infection were developed retrospectively across the nine studies and focused predominantly on symptomatic disease. Except for Quer and colleagues, each research team employed cross-validation to test their algorithm’s generalisability. Four studies randomly split their data into training and validation sets, whereas other researchers tested their algorithm on healthy and COVID-19-negative validation sets. Quer and colleagues showed that although specificity did not achieve comparably high sensitivity, With more input features, models improved in performance. Quer and colleagues showed that although the model ingesting only symptoms (AUC 0.71) performed similar to the model ingesting only wearable

| Models included in analysis | Device sensors | Manufacturer | Regulatory status | Principle of operation |
|-----------------------------|----------------|-------------|------------------|-----------------------|
| Apple Watch^2^ | Unspecified; Apple Watch Series 4 or 5 | Accelerometer, electrical heart sensor,* gyroscope, and photoplethysmography | Apple | The EU granted European conformity (CE, also known as Conformité Européenne) marking in March, 2019, for ECG app and irregular HR notifications; US FDA approved ECG app for software as a medical device, temporary approval expanded to encompass remote monitoring of heart health during the COVID-19 pandemic | The Apple Watch provides wearers with a wrist-based notification system, transmitting messages and alerts from their smartphone in real-time; it can be worn during physical activity; its battery life ranges from 1·5–18 h; in addition to supporting third-party apps, the Apple Watch includes health-focused proprietary apps; newer models (eg, the Series 6) include blood oxygen and ECG apps, in addition to the widespread irregular heart rhythm alerts |

| E4 wristband^3^ | Unspecified | Accelerometer, electrodermal activity and galvanic skin response, event mark button, infrared thermopile, internal clock, and photoplethysmography | Empatica | The EU granted CE marking to the E4 wristband, in conjunction with the complementary Aura system, in March, 2021, as a class IIa medical device intended to detect and alert users to an early respiratory infection; approval not granted yet by FDA | Lacking a hardware display, the E4 wristband enables the user to record 22 h of continuous data between device charges; it collects data through multiple sensors and transmits them to a cloud platform, storing up to 60 h of data between transfers; the device allows researchers to record biometric data of participants who are wearing the device at home or in the lab and develop their own customised apps to access participant data in real-time |

| Fitbit smartwatches and trackers^1^ | Ionic; Charge 3 and Charge 4; Inspire 2 and Inspire HR; Sense; Versa 2 and Versa 3; unspecified | Accelerometer, altimeter,* barometer,* electrical heart sensors,* GPS,* gyroscope,* orientation,* optical HR,* PurePulse 2.0 HR,* SpO2,* and skin temperature* | Fitbit | Approval not granted yet by EU or FDA | All wrist-worn Fitbit devices rely on wearable sensors to track HR, step count, and sleep stage and quality; newer smartwatch versions (eg, Sense and Versa models) also track skin temperature, SpO2, concentrations, and document potential atrial fibrillation episodes; depending on the model, Fitbit displays provide real-time measurement updates related to the wearer’s physical activity and smartphone activity; Fitbit devices can be used continuously and paired with a complementary mobile app, lasting up to 6 days between charges |

| Oura Ring^4^ | Unspecified | Accelerometer, negative temperature coefficient, photoplethysmography, and temperature | Oura | Approval not granted yet by EU or FDA | The Oura’s finger-worn design emits a physical display; designed for constant wear and is water resistant, the Oura ring has a 5–7 day battery life; the company has created an accompanying mobile app for the Oura ring; users can track their sleep, activity, and so-called readiness scores on their phone; the sleep score reflects how long the user spends in deep, rapid eye movement, and light sleep, in addition to providing personalised tips for maximising rest; the activity score considers the user’s daily steps, calories burned, and amount of time spent inactive; finally, the readiness score gives users a numeric estimate from 0 to 100 of how much their body has recovered from previous activity |

| WHOOP Strap^5^ | Unspecified | Accelerometer, capacitive touch, gyroscope, photoplethysmography, and thermometer | WHOOP | Approval not granted yet by EU or FDA | The wrist-worn WHOOP Strap collects physiological data continuously through multiple sensors; with no digital display on its hardware, the WHOOP strap’s battery lasts 4–5 days; when synced with the complementary smartphone app, the WHOOP system quantifies the user’s sleep quality, provides recommendations on how much physical exertion could be tolerated, and measures resting HR and HRV; the WHOOP app also enables users to log specific behaviours in a journal each day |

Only the named wearable devices, based on the relevant included literature, are described in the table; thus, the unnamed throat-worn patch (Lonini et al, 2021)^35 is not presented here. ECG¼electrocardiogram. FDA¼Food and Drug Administration. HR¼heart rate. HRV¼heart rate variability. SpO2¼oxygen saturation. *Model-dependent sensors.
sensor data (AUC 0.72), ingesting both symptoms and sensor data led to superior model performance (AUC 0.80). One cross-sectional study combined data from three separate devices and a self-report questionnaire to achieve an accuracy of 98.1%, compared with 82.4% when relying solely on wearable sensor data. A study enrolling patients with an influenza-like illness episode, which included COVID-19-positive individuals, showed that the symptom-based model (AUC 0.78) outperformed the wearable-based model (sensitivity 0.52, false positive rate 0.4) in distinguishing between COVID-19 cases and non-COVID-19 influenza-like illness cases. With one rare exception, the best performing models (ie, those with >90% specificity and recall of ≥80%) detected a COVID-19 infection 3–7 days after symptom onset. 

The accumulated evidence suggests a trade-off between a model’s accuracy and its ability to identify SARS-CoV-2 infection before symptom onset. Only four of the reviewed studies developed models that could detect an impending symptomatic SARS-CoV-2 infection, ranging from 14 days to the day before symptom onset. The algorithms’ ability to detect presymptomatic infection also spanned a broad range (20–88% of SARS-CoV-2 infections); however, the greater the number of days preceding symptom onset, the fewer COVID-19 cases a model could identify. For example, Mishra and colleagues detected physiological anomalies in 88% of COVID-19 cases (22 of 25 individuals with a symptom onset date) a median of 4 days before symptom onset with their model, whereas Bogu and Snyder reported detecting 56% of COVID-19 cases (14 of 25 individuals) a median of 6.94 days before symptom onset.

Heart rate, heart rate variability, respiratory rate, skin temperature, and activity levels comprised the most commonly reported physiological parameters measured by wearable devices (figure 2). We discuss the three physiological metrics that could serve as leading indicators of a SARS-CoV-2 infection, and other parameters are reviewed in the appendix (pp 15–20).

Eight articles examining data from more than three wearable devices collectively showed a positive association between SARS-CoV-2 infection and elevated heart rate. Smarr and colleagues calculated baseline physiological measurements for each Oura-wearing participant (n=50), comparing them to their mean heart rate during the first week of symptomatic infection. They found no significant difference in heart rate during illness based on participants’ self-reported symptom onset date (p=0.13), but an association with an increase in heart rate when paired with the start of device-measured temperature shifts (p=0.02). Mishra and colleagues integrated heart rate and step data from 32 Fitbit users to generate a novel heart rate over steps feature. Their analysis revealed that, among 25 individuals with discernible changes in their physiological parameters around symptom onset, heart rate increased by a median of 7 beats per min. Using a subset of Mishra and colleagues’ data, Bogu and Snyder developed an algorithm to detect anomalies in resting heart rate around the time of a potential SARS-CoV-2 infection and reported that COVID-19-positive individuals had more recorded hours of abnormal heart rate during the infectious period than healthy peers or those who were ill from a cause other than COVID-19. Although heart rate anomalies could help alert a wearable device user to an impending infection, research suggests changes in heart rate alone cannot differentiate SARS-CoV-2 infection from other influenza-like illnesses. Shapiro and colleagues showed that both patients with COVID-19 and patients with influenza had elevated heart...
rate following self-reported symptom onset. In their device-agnostic studies, Quer and colleagues\(^\text{34}\) and Cleary and colleagues\(^\text{35}\) found no relative difference between elevated heart rate in COVID-19-negative cohorts and COVID-19-positive cohorts (p=0.33 and p=0.18).\(^\text{34}\) Furthermore, the same machine learning model ingesting a heart-rate-derived feature could not discriminate well between COVID-19-positive individuals and COVID-19-negative individuals (AUC 0.52 and 0.63).\(^\text{34}\) In another study,\(^\text{39}\) even variability in heart rate before and after activity remained similar, regardless of health status. Converging evidence suggests intrapersonal heart rate might increase following a SARS-CoV-2 infection, but it cannot serve as the sole discriminating factor.

Three of the four studies examining SARS-CoV-2 infection’s effect on respiratory rate found that it increased around symptom onset.\(^\text{34}\) In one study,\(^\text{38}\) SARS-CoV-2-positive Oura users had higher respiratory rates during the early symptomatic period than during the pre-illness baseline (p=0.002). Training a convolutional neural network on physiological data from 1257 Fitbit wearers, Natarajan and colleagues\(^\text{35}\) reported that, during a SARS-CoV-2 infection, respiratory rate deviated from its baseline value more than other parameters. In contrast, Miller and colleagues\(^\text{39}\) did not identify respiratory rate as a leading indicator of a potential SARS-CoV-2 infection in their examination of 271 WHOOP strap users who reported COVID-19 symptoms; compared with other physiological parameters, respiratory rate had the lowest coefficient of intraindividual variance over time, regardless of whether the patient was healthy or ill on a given day.

Whereas other articles considered deviations in respiratory rate during a SARS-CoV-2 infection compared with a previous baseline period, Lonini and colleagues\(^\text{39}\) examined physiological changes occurring on the same day before and after a given activity. The researchers equipped 15 participants with SARS-CoV-2 infection and 14 healthy participants with an unnamed wearable device. Patients positive for SARS-CoV-2 had similar respiratory rate variability in response to exercise compared with healthy peers (p=0.095), despite a higher baseline value. Cohort demographic differences, however, limit the generalisability of their findings, as most COVID-19 cases had a comorbidity that could have affected their baseline respiratory rate (eg, asthma).\(^\text{39}\)

Although fever was one of the first COVID-19 symptoms identified by WHO,\(^\text{2}\) of the studies that measured skin temperature, only Smarr and colleagues\(^\text{30}\) focused on assessing deviations in this physiological parameter. They compared Oura users’ baseline skin temperature to the period following self-reported COVID-19 symptom onset. Statistical analysis revealed an increase in temperature during a symptomatic SARS-CoV-2 infection (p=0.024), with 76% (38 of 50) of participants registering an increase in temperature in the days preceding symptom onset.

We evaluated risk of bias on the basis of the National Institutes of Health’s Quality Assessment Tool for Observational Cohort and Cross-sectional Studies.\(^\text{29}\) We provide a study-by-study breakdown and detailed descriptions of individual biases in the appendix (pp 21–24). In general, most studies presented a moderate risk of bias; the definition, size, self-reporting of diagnosis, and demographics of the study populations represented a major source of potential bias. Several articles did not clearly define the study population (eg, age, comorbidities, and nationality).\(^\text{30,34,35,37,39}\) Three studies also had small samples (total analysed sample n=1500), despite starting from very large recruited populations (>30 000 individuals).\(^\text{30,34,35}\) Some researchers attempted to address the restricted sample size and class imbalance (ie, the number of participants who were positive and negative for COVID-19, or positive and negative days, depending on the type of observation that was analysed) in their algorithms by upsampling infection days,\(^\text{30}\) implementing bootstrapping with replacement,\(^\text{39}\) or generating a synthetic training dataset.\(^\text{36}\) Moreover, most studies identified SARS-CoV-2 infection through participant self-report,\(^\text{39}\) which is overly reliant on subjective data and potentially misses asymptomatic cases. Confounding represented a source of bias faced by many studies, given their restricted adjustment for major demographic factors. Furthermore, many pre-existing comorbidities (eg, body-mass index)\(^\text{35}\) shown to affect COVID-19 vulnerability and severity\(^\text{39}\) were rarely ingested by the algorithms.\(^\text{30,34,35,39}\)

In addition to the articles detailing completed research, 12 study protocols met inclusion criteria.\(^\text{41–52}\) The protocols are investigating numerous wearable devices, ranging from a repurposed fertility tracking bracelet\(^\text{43,49}\) to a wearable device supplemented by a sensor placed under the participant’s mattress.\(^\text{44}\) These studies aimed to assess changes in physiological parameters commonly examined by the other studies we included in our analysis, including heart rate, heart rate variability, and temperature. At least one protocol intends to examine a previously unreported parameter (ie, blood pulse wave).\(^\text{30}\) Unlike the completed studies, all protocols propose prospective studies, including two RCTs.\(^\text{43,49}\) Two protocols plan to include healthy control groups.\(^\text{41–51}\)

**Discussion**

This systematic review examined 12 publications and preprints and 12 study protocols related to wearable devices’ ability to detect a potential SARS-CoV-2 infection. We observed large variability in device type, physiological parameters analysed, and the operationalisation of diagnostic accuracy across models. Some authors relied on statistical analysis to detect differences between or within participants, whereas others used machine learning algorithms. Accordingly, models varied in their feature specification and performance.
At present, the overall body of evidence regarding the use of wearable devices to detect COVID-19 shows promising, albeit early stage, findings. Most studies drew on retrospective data, had small sample sizes, and did not examine physiological differences from other influenza-like illnesses. Although some studies used PCR testing to confirm SARS-CoV-2 infection, this practice was not universally deployed, potentially introducing diagnostic biases and restricting the comparability of studies to each other. Only three of the included studies explicitly reported using PCR testing—either a weekly PCR test, a per occurrence self-reported PCR test, or a one-time PCR test upon hospital arrival. The fact that two of those studies were conducted solely or partly on health-care professionals shows that this type of population might have had easier access to PCR testing, because of job requirements during the first wave of the COVID-19 pandemic, than the general population. Each study had a different design (ie, prospective, retrospective, and cross-sectional) and investigated the ability of different devices (ie, Apple-Watch, Fitbit, and Empatica E4) to detect deviations in physiological parameters associated with a SARS-CoV-2 infection. Of note, the only included studies that present findings for participants who are infected but asymptomatic, report the use of PCR testing as the reference test. Their relatively small sample sizes and prospective and cross-sectional designs could have made it feasible to require PCR testing during data collection. Also, two studies that used PCR tests to determine infection and that aimed to classify the current infection status of participants by developing neural networks, achieved high accuracy (98.1%- and 77.0%-) and specificity (99.0%-). However, this performance cannot necessarily be attributed to reliance on the gold-standard PCR tests as an infection marker, as multiple other differences in the specifications and inputs to their models could have influenced their capabilities for detecting a SARS-CoV-2 infection. For example, Natarajan and colleagues enrolled a large sample (n=1257) of symptomatic Fitbit users and examined the classification of a given day for each individual as healthy or ill based on preceding physiological data of heart rate, heart rate variability, and respiratory rate, as well as demographic characteristics. In contrast, Hassantabar and colleagues analysed a much smaller sample (n=87) of healthy (negative PCR test) and symptomatic and asymptomatic patients infected with SARS-CoV-2 (positive PCR test) by monitoring data for up to 1 h from multiple devices measuring galvanic skin response, SpO2, and blood pressure, and collecting questionnaire data on demographics, symptoms, and comorbidities. Despite disparate methods and differences in follow-up time, both studies validated high-performing machine learning algorithms for diagnosing a SARS-CoV-2 infection. Less than half of the studies included a control group of healthy participants, which further limits generalisability. Findings from several studies on changes in physiological parameters might also appear contradictory at first glance. However, discrepancies in the direction or magnitude of change could be attributable to the brand or model of a given device. For example, both Miller and colleagues and Natarajan and colleagues analysed data from a wrist-worn device and arrived at differing conclusions about the effect of SARS-CoV-2 infection on respiratory rate. However, the sensors in the specific hardware or the underlying data extraction techniques for interpreting raw sensor data could vary substantially between the WHOOP strap studied by Miller and colleagues and the Fitbit bracelet studied by Natarajan and colleagues. Both wearable devices have a photoplethysmography sensor; however, differences in sampling rates for the sensors could explain variations in interbeat intervals and derived respiratory rates. Device-agnostic studies, pooling data from multiple device models and brands, might seem to directly address these discrepancies through uniform data processing and algorithm development; yet similar concerns could nevertheless render their results difficult to interpret. Recruiting Fitbit users and collecting data from participants’ Apple HealthKits and Google-based devices, Quer and colleagues did not correct for potential confounding biases related to the different wearables. Their finding of no changes in heart rate on the basis of SARS-CoV-2 infection status could have derived from how each wearable device measures and processes its raw physiological data. Subsequent device-agnostic studies could further clarify the relationship between seemingly discrepant findings by conducting a head-to-head comparison and determining whether a model’s performance varied by device type.

Beyond biases introduced by differences in the studied wearable devices and their associated sensors, the included articles also lacked standardisation in their algorithm development and reporting of performance metrics. After receiving the raw physiological data from a wearable device, researchers make decisions regarding the signal’s preprocessing and cleaning (eg, normalising or transforming data before model training). Additionally, researchers must choose which optimiser to use when training their model; different optimiser selections can affect model fit and performance. The included studies used more than five types of optimisers in training their respective models. Best practice in machine learning also suggests having a separate test or validation dataset from the training dataset; although two studies did not include any test set, seven articles varied greatly in their approach to validating their machine learning algorithm, and the remaining three studies used statistical analyses rather than machine learning methods. Miller and colleagues tested their model on a dataset of participants derived from the same population as their training data, although their data were recorded during a different time period; in contrast,
Hassantabar and colleagues relied on a categorically different population (ie, healthy controls) to validate their algorithm. How each research team chose their test or validation sets inherently influenced their algorithm’s performance. Showing how the algorithm development process affects performance, Nestor and colleagues reported their model’s sensitivity, specificity, and other metrics if held to the same evaluation schemes as other authors. Their best performing model achieved a higher specificity and sensitivity compared with Miller and colleagues. Recently, the scientific community has recognised the need for improved standardisation in algorithm development and performance metric reporting, particularly as they relate to health outcomes; multiple publications have called for clinical trials to use machine learning techniques to show their data input process, handling of missing or poor-quality data, and outcomes. Although aligned with current practices in machine learning, the varied approaches to algorithm development documented across the included studies could have introduced bias in the findings.

Except for two papers, the examined studies also did not consider how physiological parameters might differ between symptomatic and asymptomatic SARS-CoV-2 infections. Most authors trained their algorithms exclusively on symptomatic infections. Identifying asymptomatic infections and building a corresponding model requires testing participants repeatedly for SARS-CoV-2 antibodies; a procedure not done by any of the studies included in our systematic review. Using specialised anomaly-detection algorithms, researchers could train a machine learning model to recognise intrapersonal deviations in physiological parameters during the period between baseline seronegativity and known seroconversion. This model could retrospectively identify the timing of a previous SARS-CoV-2 infection and be applied prospectively to determine real-time asymptomatic—but nevertheless still transmissible—infections. A single protocol, identified by our literature search, was that none of the models detecting SARS-CoV-2 infection on the basis of physiological parameters were tested or validated in real-time; although one study tested their online algorithm retrospectively. An algorithm-informed real-time indicator that ingests wearable sensor data could enable individuals to make behavioural changes, such as seeking a SARS-CoV-2 test early and self-isolating. Another study simulated real-world deployment, warning that the shifting prevalence of COVID-19 could cause substantial overestimation of model performance. All identified protocols, however, follow a prospective design, with three protocols aiming to assess an algorithm-driven alert system for health-care professionals or the wearable device users. Future research plans thus show the need and desire to address this gap.

This systematic review also highlights the disproportionate representation of wrist-worn devices in research surrounding SARS-CoV-2 detection, restricting the results’ potential generalisability. Four of the five named devices were smartwatches or wrist-worn straps, whereas only two of the 12 included articles (17%) studied physiological changes related to a SARS-CoV-2 infection with other types of wearable devices (eg, a smart ring). We designed our search strategy to minimise potential bias in method of wearable device by including generic terms (eg, smart sensing technology) and searching specifically for non-wrist-worn devices (eg, skin patch and smart glasses; see appendix pp 3–5 for a full list of search terms). Nevertheless, the literature was skewed heavily towards wrist-based wearable devices. An inherent limitation resulting from this disproportionate representation, differences in sensor types, size, and placement could lead to variations in their measurements and accuracy. Although it is beyond the scope of this systematic review to dissect the engineering and design principles varying across wearable devices, we acknowledge the preponderance of wrist-based wearable devices in the summarised literature might unduly influence our findings and conclusions. More studies focused on non-wrist-worn devices will be needed to disentangle how device type influences algorithm performance and which physiological parameters change in relation to SARS-CoV-2 infection. Five protocols identified by our search include non-wrist-worn sensor components, and the results of these studies will contribute much-needed data to this body of evidence.

Despite constituting key features affecting participant compliance and overall adoptability, the wearable and perceived ease of use for each studied wearable device were not discussed in any of the included studies. Additionally, comparing usability across devices is difficult because of differences in study design. For example, snapshot studies recruiting small samples for a short period of time placed a small burden on participants in terms of time and effort; a participant might be more likely to tolerate an uncomfortable device if they need only to wear it for a few hours compared with a study lasting several months. Conversely, studies using stand-alone consumer wearables relied on users to opt-in to their clinical trials recruiting small samples placed a small burden on participants might have felt more comfortable with the device they already own compared to study participants who were given the
device as study material.31-33 Many of the included studies required a minimum time or days of use of the wearable device as a prerequisite for a participant to be included in the analysed sample.30-33,35-38 However, there were differences in how long users had to wear the device each day between studies with extended timelines. For example, some studies30,35 developed their models using only night data, although the devices were designed to be worn throughout the day. Consequently, their findings suggest individuals need only wear the device while at rest to detect a SARS-CoV-2 infection. Usability and perceived ease-of-use would probably be affected by how often a user has to wear the device, in addition to its baseline comfort. Finally, some researchers observed that participants occasionally did not use the wearable device when symptomatic,19 indicating participant’s health could interact with the device’s overall wearability, affect data collection, and subsequently the underlying model’s ability to detect an infection. This research team also argued that devices requiring daily charging are expected to have more missing data, showing that this feature potentially affects the data quality of the study.19 Although another research team analysed all participants in the cohort and used machine learning models that can handle implicitly missing data for this purpose, they report a drop in performance if an individual had not worn the device for a week.12 The fact that several of the included study protocols aim to assess the feasibility of wearing a device for a specific amount of time is encouraging.6,44,45 Various future studies intend to establish individuals’ comfort in following the necessary compliance schedule to maximise the usefulness of a SARS-CoV-2 detection algorithm. One trial designed for the specific intensive settings of a 14-day quarantine will instruct participants to wear the device all the time except for when showering and charging the device.19 Similarly, another planned study will ask participants to wear the device all the time outside of work (health-care workers) for 30 days, but recognises in their protocol that there is a small chance of discomfort or skin abrasion from the prolonged use of the wristband without following appropriate hygiene practices.22 One trial will aim to balance the prolonged follow-up of the participants with asking them to wear the monitoring bracelet only at night, so that the skin can breathe and dry during the day.41 Although we cannot comment on the wearability or ease-of-use of the studied devices, future synthesis of these factors for each device should be feasible given the clinical trials underway.

Finally, sample selection and participant demographics limit the models’ generalisability across populations. For example, most studies, which were done solely or largely in the USA, had little racial diversity, despite COVID-19 disproportionately affecting Black and Hispanic communities in the USA.29 Wearable devices have previously shown variable performance across differing skin tones; consequently, if research does not explicitly include these populations, diagnostic accuracy and potential for public good within vulnerable communities remains limited.29 Similarly, although some models attempted to take into account sex-based variance,6,45 none of the machine learning algorithms considered how physiological parameters change across the menstrual cycle.19-41 Future researchers should consider and fine-tune their algorithms to adjust for sex-based differences, thereby reducing the likelihood that a postovulatory shift in temperature, for example, would be erroneously labelled as COVID-19.

Despite these limitations, the reviewed studies provide valuable insights for future research on common wearable-measured physiological parameters. Eight articles showed an increase in heart rate associated with a SARS-CoV-2 infection,29,32-34,36-40 in line with population-level heart rate data associated with influenza.29 Similarly, changes in skin temperature and activity frequency provide encouraging results for emerging wearable devices equipped with a temperature sensor and an accelerometer. Notably, behavioural changes, such as receiving a SARS-CoV-2 test result after symptom onset could result in an overestimation of the performance of models based on activity frequency, limiting the use of these data samples.31 In addition, establishing a conclusive COVID-19-related pattern in respiratory rate and heart rate variability requires additional replications to disentangle contradicting or inconclusive initial findings. Other features, such as coughing patterns from mechaanoacoustic sensors could be used to decipher further trends relevant to a SARS-CoV-2 infection,59 although this possibility remains to be shown. Several experimental papers also used this approach,29 although their content did not fit the inclusion criteria for this systematic review.

This study represents a comprehensive search of multiple databases and literature to date; we included multiple synonyms for each primary term, tailored the search terms to each database, manually screened reference lists, and actively tried to mitigate any missing data. Owing to the rapid pace of COVID-19 research, we also sought out preprint sources. Despite these strengths, although we did not restrict language in the databases we used, we did not search databases publishing only non-English publications; thus we might have overlooked relevant literature published in another language. Additionally, we identified only studies captured according to our search terms, in specific medical and research databases. Despite our efforts, we might have missed some relevant studies which have not yet been published in peer-reviewed journals or as preprints (eg, findings reported in news articles or company press releases). We sought to mitigate this potential limitation by including a supplementary search for study protocols, which could highlight ongoing clinical trials and potential future publications relevant to our research question.
Conclusion

Adequately containing the COVID-19 pandemic requires rapid identification of individuals who are infectious. Although wearable devices could help, this systematic review highlights the need for well designed and controlled studies to robustly identify if wearables can accurately detect SARS-CoV-2 infection before symptom onset or in asymptomatic individuals in comparison to the current gold-standard diagnostic method. Future studies should additionally consider how inherent differences in wearable sensor methods, raw data processing, and algorithm development contribute to the detection of infection-associated deviations in physiological measurements and how to address sources of bias.

Contributors

All authors had full access to the data, including statistical reports and tables, in the study. MM, BMG, and GSD take responsibility for the integrity of the data and the accuracy of the data analysis. MM, BMG, DEG, PS, MC, and GSD contributed to the study concept and design. MM, BMG, AS, VK, and GSD contributed to the acquisition, analysis, and interpretation of data and provided administrative, technical, and material support. MM, BMG, and GSD contributed to drafting the manuscript. All authors contributed to a critical revision of the manuscript for important intellectual content. DEG, PS, and MC obtained funding. GSD was responsible for study supervision.

Declaration of interests

MM, BMG, VK, BF, DV, DEG, MC, and GSD received grants from Innovative Medicines Initiative 2 Joint Undertaking (number 101005177), during the conduct of the study. BMG reports consulting fees and employment from Ava Science, support for attending meetings and travel from Ava Aktiengesellschaft (AG), a patent application from Ava AG (P24892CH100) filed with the Swiss Federal Institute of Intellectual Property for System and Method for Pre-Symptomatic and/or Asymptomatic Detection of a Human Viral or Bacterial Infection based on pilot data from the COVID-RED clinical study, and consultancy for Falcon Health and Therab Medical, outside the submitted work. VK reports employment from Ava Science and Ava AG, during the conduct of this study. TB, BF, DV, and DEG report employment from Julius Clinical Research, during the conduct of the study. MC reports employment from Ava AG during the conduct of the study. GSD reports a grant from Health Holland, outside the submitted work. All other authors declare no competing interests.

Data sharing

Template data collection forms and the completed data extraction table are included in the appendix (pp 26–40).

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The Data sharing section describes the data collection forms and the completed data extraction table, which are included in the appendix.

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The References section includes a list of publications with hyperlinks, indicating that they can be accessed online. The references are cited in the text, and the page numbers are provided for easy reference.
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