

# Classification of Patient Recovery From COVID-19 Symptoms Using Consumer Wearables and Machine Learning

Jared Leitner<sup>1</sup>, Alexander Behnke, Po-Han Chiang<sup>1</sup>, *Student Member, IEEE*, Michele Ritter, Marlene Millen, and Sujit Dey, *Fellow, IEEE*

**Abstract**—Current remote monitoring of COVID-19 patients relies on manual symptom reporting, which is highly dependent on patient compliance. In this research, we present a machine learning (ML)-based remote monitoring method to estimate patient recovery from COVID-19 symptoms using automatically collected wearable device data, instead of relying on manually collected symptom data. We deploy our remote monitoring system, namely eCOVID, in two COVID-19 telemedicine clinics. Our system utilizes a Garmin wearable and symptom tracker mobile app for data collection. The data consists of vitals, lifestyle, and symptom information which is fused into an online report for clinicians to review. Symptom data collected via our mobile app is used to label the recovery status of each patient daily. We propose a ML-based binary patient recovery classifier which uses wearable data to estimate whether a patient has recovered from COVID-19 symptoms. We evaluate our method using leave-one-subject-out (LOSO) cross-validation, and find that Random Forest (RF) is the top performing model. Our method achieves an F1-score of 0.88 when applying our RF-based model personalization technique using weighted bootstrap aggregation. Our results demonstrate that ML-assisted remote monitoring using automatically collected wearable data can supplement or be used in place of manual daily symptom tracking which relies on patient compliance.

**Index Terms**—Machine learning, wearables, remote patient monitoring, COVID-19.

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Jared Leitner, Alexander Behnke, and Po-Han Chiang are with the Electrical and Computer Engineering Department, University of California San Diego, La Jolla, CA 92093-0021 USA (e-mail: jleitne@eng.ucsd.edu; abehnke@eng.ucsd.edu; pochang@ucsd.edu).

Michele Ritter and Marlene Millen are with UCSD Health, San Diego, CA 92103 USA (e-mail: mlritter@health.ucsd.edu; mmillen@health.ucsd.edu).

Sujit Dey is with the Electrical and Computer Engineering Department and Head of the Mobile Systems Design Laboratory, University of California San Diego, La Jolla, CA 92093-0021 USA (e-mail: dey@ece.ucsd.edu).

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## I. INTRODUCTION

AROUND the world, healthcare systems have been overwhelmed by the high numbers of COVID-19 cases, which has surpassed 437 million as of March 2, 2022 according to the World Health Organization (WHO) [1]. In the US, there were approximately 4.5 million COVID-19 hospitalizations between August 1, 2020 and February 28, 2022, according to the Center for Disease Control and Prevention (CDC) [2]. While this is a daunting number of hospitalizations, there have been approximately 80 million cases in the US [3], meaning most cases involve ambulatory patients being treated from home. This is an unprecedented number of patients needing care in their home and many are not being monitored in any way by medical personnel.

In order to combat this pandemic and provide more optimal care at scale, hospitals are changing the way in which healthcare is delivered. At the center of this changing landscape is a shift towards remote, continuous, and automated delivery of healthcare. This shift can lead to significant improvement in and scalability of at-home patient care for COVID-19, while at the same time enabling significant savings in human and equipment resources. Current remote monitoring for COVID-19 patients relies on manual symptom reporting, which is highly dependent on patient compliance. In this study, we demonstrate that data automatically collected from wearable devices together with machine learning (ML)-assisted diagnosis can enhance the efficiency and increase the scalability of remote monitoring for COVID-19 patients.

Wearable devices are one of the enabling technologies making this shift in healthcare delivery possible [4], [5], [6], [7]. Consumer wearables, such as Apple Watch, Fitbit, and Samsung Galaxy Watch, remotely collect a great amount of lifestyle and vitals data in high granularity and continuity. There is great opportunity for ML to assist in remote monitoring due to the large amount of data that is collected. Since it is not possible for doctors to manually review all remotely collected data [8], ML has the potential to provide automated insights into the health status of patients and significantly increase the scalability of remote patient care. This is especially helpful during a pandemic, where in-person interaction and monitoring may pose risks to healthcare workers and other patients. In addition, ML-assisted monitoring can provide patients with insights regarding their own progression, helping to keep them engaged and informed about their health.

Current research on using wearables and machine learning to combat COVID-19 is primarily focused on early detection of infection. The authors in [12], [13], [14], [15], [16] have

demonstrated that it is possible to detect deviations in health data before significant symptoms arise. Using Fitbit devices, the researchers in [12] found that 26 out of 32 (81%) infected patients in their cohort had alterations in their heart rate, number of daily steps, or time asleep before becoming symptomatic. The authors in [15] used respiration rate, heart rate, and heart rate variability data collected from their wearable devices and proposed a deep learning method to estimate infection before the onset of symptoms. Early detection will enable individuals to quarantine earlier, helping reduce the spread of the virus. These studies demonstrate that wearable device data can provide actionable insights into the conditions of patients.

In this research, we propose a novel approach to estimate patient recovery from COVID-19 symptoms using automatically collected device data and machine learning. We partnered with the UCSD Health and Neighborhood Healthcare COVID-19 telemedicine clinics in order to carry out this research. Our remote monitoring system utilizes a Garmin wearable and symptom tracker mobile app for data collection and fuses this data into an online report for clinicians to review. We propose a novel labelling logic for patient recovery from COVID-19 symptoms using the symptom tracker data. The labelling logic was developed in collaboration with UCSD Health doctors and the details are defined in Section III-B. Using this data, we train a patient recovery classifier which uses wearable data to estimate whether a patient has recovered from COVID-19 symptoms. We evaluate our method according to leave-one-subject-out (LOSO) CV to replicate the clinically relevant use case scenario in which a newly infected patient will not have data for model training. We compare the performance of different ML models and find that Random Forest (RF) is the top performing model. We propose a RF-based personalization technique in order to improve model performance. This technique utilizes the RF's weighted bootstrap aggregation algorithm in order to tune the model to each patient. The details are presented in Section III-D. Finally, we conduct Shapley Value analysis to inspect which device features have the greatest impact on classification. This analysis provides an interpretation of what the model has learned, which is especially important for medical applications. Our contributions are as follows:

- We deploy a remote patient monitoring system in two COVID-19 telemedicine clinics. The system consists of a wearable device, symptom tracker mobile app, and online dashboard which collects and analyzes vitals, lifestyle, and symptoms data. The estimated recovery status of each patient using our ML approach is displayed on the dashboard for clinicians to review.
- We propose a patient recovery classifier which uses wearable data to estimate whether a patient has recovered from COVID-19 symptoms. This ML tool can provide doctors with automated insights into the recovery status of their infected patients and bypass the need for manual daily symptom tracking.
- We carry out LOSO CV to mirror the clinically relevant use-case scenario and propose a RF-based personalization technique that improves model performance by tuning the model to each patient via weighted bootstrap aggregation.

The rest of the paper is organized as follows. In Section II, we investigate related works that utilize machine learning for COVID-19 diagnosis. In Section III, our remote monitoring

system and data acquisition are presented. We then detail the proposed labelling logic and RF-based personalization technique for patient recovery classification. In Section IV, the performance of our proposed ML method is evaluated. In addition, we carry out top feature analysis based on Shapley Values and provide a discussion on research challenges. Finally, we conclude the paper in Section V.

## II. RELATED WORK

In this section, we present related research which is grouped into the follow categories: COVID-19 symptom tracking, early diagnosis, and recovery detection.

### A. COVID-19 Symptom Tracking

The researchers in [9] utilize a smartphone-based app to collect symptom data from patients. In the app, patients also recorded when they had tested either negative or positive for COVID-19 infection. They propose a logistic regression model that combines the reported symptoms in order to predict COVID-19 infection. A combination of loss of smell and taste, fatigue, persistent cough, and loss of appetite resulted in the best model, which achieved a sensitivity and specificity of 0.65 and 0.78, respectively. The authors in [10] also used a mobile app for collecting symptoms data and COVID-19 test results. They trained a logistic regression model to predict COVID-19 infection based on self-reported symptoms, and calculated the odds ratio for each symptom in order to understand which symptoms were the strongest predictors. Chills, fever, loss of smell, nausea, and shortness of breath were the top five strongest predictors of COVID-19 infection. Participants in their cohort with a positive test result experienced 5.6 symptoms on average. In [11], the researchers trained a gradient-boosting machine to predict COVID-19 infection based on 8 features: cough, fever, sore throat, shortness of breath, headache, age, sex, and known contact with an individual confirmed to have COVID-19. Their approach achieved a sensitivity and specificity of 0.86 and 0.79, respectively. Fever and cough were the top 2 features with the greatest impact on the model's prediction. These past works demonstrate that self-reported symptoms can be effectively used to predict COVID-19 infection. However, these approaches rely on patient compliance with manual symptom tracking. In contrast, wearable devices can passively collect data that is relevant to COVID-19 infection. In addition, wearable data can be predictive of COVID-19 infection prior to symptom onset.

### B. Early Diagnosis of COVID-19

The authors in [12] use data collected from wearable devices for the early detection of COVID-19 infection. They propose an anomaly detection technique based on two parameters: 1. Resting heart rate (RHR), 2. Heart rate over steps (HROS). HROS was calculated by dividing heart rate by steps data at each hourly interval. They report that significant deviations in these parameters relative to the individual baseline can indicate COVID-19 infection. They utilize Gaussian density estimation to classify anomalies in the dataset. Their results show that 63% of COVID-19 cases in their cohort could have been detected before symptom onset. The researchers in [13] also utilize deviations from RHR to classify a patient as infected. They propose a deterministic finite state machine which triggers an alert when a

patient's overnight RHR increases above the median of previous overnight RHRs by an empirically determined threshold. Their system generated alerts for 80% of the infected individuals prior to symptoms, however, many of the alert-generating events were not associated with COVID-19 and instead attributed to other events, such as poor sleep, stress, alcohol consumption, intense exercise, or travel. While these studies demonstrate that deviations in physiological and activity data measured by wearable devices can be used for early detection of COVID-19, they only utilize a subset of possible device features (RHR and steps) and do not investigate ML-based approaches which are well suited to handle larger feature sets. Furthermore, they do not investigate if wearable device data can be used to monitor patient recovery from COVID-19.

The researchers in [14] trained a logistic regression model to differentiate COVID-19 positive vs. negative cases in symptomatic individuals based on symptoms and wearable device data. Baseline device data was calculated as the median of the data from 21 to 7 days before the onset of symptoms. They show an increase in model performance when including device data (RHR, sleep duration and step count) in addition to symptoms data as part of the feature set. The authors in [15] trained a convolutional neural network to predict illness given health metrics for that day and the preceding 4 days. These metrics included the mean respiration rate (RR) during sleep, mean heart rate during sleep, the root mean square of successive differences (RMSSD) of the nocturnal RR series and the Shannon entropy of the nocturnal RR series. They organize each data sample into  $5 \times 4$  matrix and resize each matrix into a  $28 \times 28$  image as the input to the network. Their method achieved a sensitivity and specificity of 51% and 90%, respectively. In [16], the researchers presented a gradient-boosting model based on decision trees to detect COVID-19 infection. Their approach achieved a sensitivity and specificity of 71% and 67%, respectively, when only using device features as input to the model. They grouped the device features into activity, sleep, and heart rate categories, and found that activity related features had the greatest impact on the model's prediction, followed by sleep and heart rate-related features. These works demonstrate the ability of ML models to learn meaningful relationships between wearable device features and the onset of COVID-19 infection.

### C. Recovery Detection From COVID-19

The research presented in [9], [10], [11], [12], [13], [14], [15], [16] focused on predicting COVID-19 infection using self-reported symptoms or wearable device data. In contrast to these works, the objective of our research is to estimate recovery from COVID-19 symptoms using wearable device data. The researchers in [17], [18] present different approaches to estimate recovery from COVID-19 infection based on symptoms and demographic data. The authors train a support vector machine [17] and decision tree classifier [18] to estimate patient recovery based on symptoms, demographic, and travel-related features. In [17], the authors found that most of the patients who could not recover experienced a fever, cough, and fatigue. In [18], the authors extended their model to predict the number of days needed to recover from infection. Their model predicted a minimum of 5 days and a maximum of 35 days for COVID-19 patients to recover. Both approaches presented in [17], [18] rely on symptoms data and do not investigate the use of wearable device data for patient recovery estimation. We did not find any previous

TABLE I  
COHORT STATISTICS (N = 30)

	UCSD Health	Neighborhood Health
Total	23	7
# Men	11	3
# Women	12	4
Age (years, mean $\pm$ SD)	44.5 $\pm$ 13.1	31.6 $\pm$ 13.5

research that investigates whether wearable device data can be used to estimate patient recovery from COVID-19. This aligns with the observations of the authors in [19] who provide a review on the rise of wearables during the COVID-19 pandemic. None of the works presented in their review are focused on estimating patient recovery from COVID-19 symptoms. This motivates us to develop our own labeling logic for patient recovery in direct consultation with UCSD Health COVID-19 telemedicine doctors. In addition, the dataset we collect consists of a rich feature set spanning activity, sleep, stress, heart rate and SpO<sub>2</sub> data. Our paper provides novel insights into which lifestyle and physiological signals are associated with patient recovery from COVID-19 symptoms.

## III. METHOD

In this section, we first detail our study cohort and the proposed remote patient monitoring and reporting system. We then present the ML task of patient recovery classification and discuss its application. Finally, we describe the data preprocessing, the RF model, and our proposed personalization technique.

### A. Clinical Study Cohort and eCOVID System

Our IRB-approved clinical study (protocol #181405) was in collaboration with UC San Diego Health and Neighborhood Healthcare, with patient enrollment, onboarding and management conducted by the Altman Clinical & Translational Research Institute at UC San Diego. The study was conducted starting in May 2020. Patients who tested positive for COVID-19 at each location were referred to our study coordinator. Eligible patients were required to be over 18 years old and stable for monitoring in an ambulatory setting, as determined by healthcare personnel at the point of care when testing was initially ordered. The characteristics of the included cohort are shown in Table I. Subjects digitally consented using our symptom tracker mobile app, and those who did were provided a Garmin Vivosmart4 wearable device [20] to collect their lifestyle and vitals data for the study duration of up to 3 months. One of the deciding factors in using this device for this study is its ability to measure blood oxygen saturation (SpO<sub>2</sub>). Based on the findings of [21] and our discussion with UCSD Health doctors, SpO<sub>2</sub> is a critical metric in determining the condition of a COVID-19 infected patient. Fig. 1 displays the overall architecture of our remote monitoring system, namely eCOVID. The system consists of a symptom tracker mobile app, developed using the Touchwork platform, and the Garmin device. The daily questions in the symptom tracker app were developed in collaboration with doctors at the UCSD Health COVID-19 telemedicine clinic and are detailed in Table II. The vitals and lifestyle data collected by

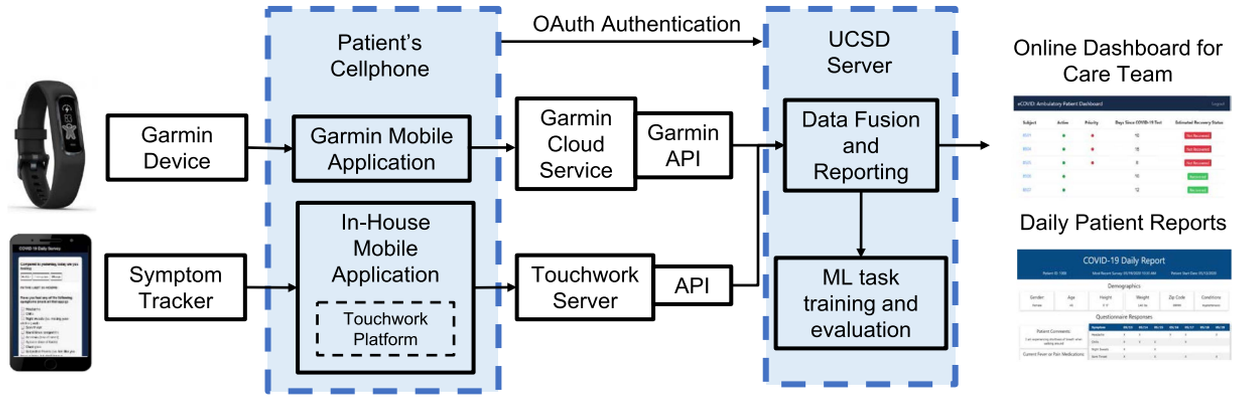


Fig. 1. eCOVID remote monitoring and reporting system architecture.

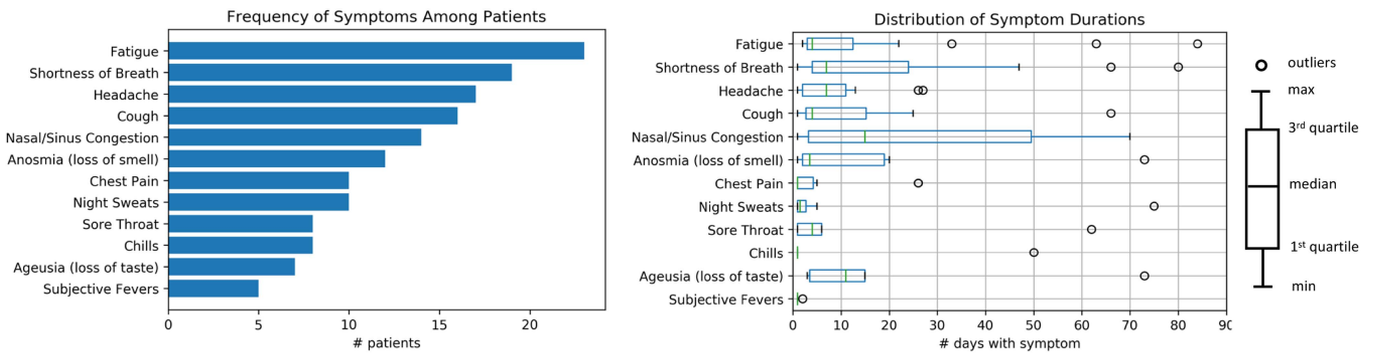


Fig. 2. The left plot displays the number of patients who reported at least 1 day of the symptom. The right plot displays the distribution of the number of days each symptom was reported per patient. Only patients who reported the symptom are included in this distribution.

TABLE II  
DAILY QUESTIONS IN SYMPTOM TRACKER APP

Questions (Answers)
1. How do you feel compared to yesterday? (Better, Same, Worse)
2. Have you had any of the following symptoms? (Headache, Chills, Night sweats, Sore throat, Nasal/sinus congestion, Anosmia, Ageusia, Chest pain, Subjective fevers)
3. How would you rate your fatigue? (0-5)
4. How would you rate your cough? (0-5)
5. How would you rate any shortness of breath? (0-5)
6. Are you able to drink & eat? (Yes, Somewhat, Little, Minimal)
7. What fever/pain medications have you taken?
8. What cough/breathing medications have you taken?

the Vivosmart4 wearable are detailed in Section III-C. Data was collected remotely through Garmin’s application programming interface (API) [22].

Fig. 2 details the distribution of symptoms among patients and describes how long each symptom lasted. For fatigue, shortness of breath and cough, we marked the symptom as present if the patient reported a severity score of 2 or greater. The bar graph in Fig. 2 displays the number of patients that experienced each symptom. Fatigue, shortness of breath and headache were the

3 most common symptoms with 23 (77%), 19 (63%) and 17 (57%) patients reporting these symptoms, respectively. Chills, ageusia and subjective fevers were the 3 least common symptoms with 8 (27%), 7 (23%) and 5 (17%) patients reporting these symptoms, respectively. The box plot in Fig. 2 details how long each symptom was reported by patients. Only patients who reported the symptom are included in this analysis. Based on the median number of days, nasal/sinus congestion lingered the longest with a median of 15 days followed by ageusia with a median of 11 days. Although ageusia was only reported by 7 patients, the symptom lingered for a longer time compared to other symptoms. Subjective fevers, chills and chest pain were reported for the shortest period each having a median of 1 day. Patients completed the daily symptom tracker an average of 73% of days enrolled in the study. They wore the Garmin device an average of 90% of days enrolled in the study. This indicates that patient compliance with wearing the device was 17% greater than compliance with answering the daily symptom tracker. This statistic demonstrates the higher efficiency of wearable device data for remote monitoring and helps motivate our proposed ML task for patient recovery classification based on automatically collected device data, as opposed to relying on manually entered symptom data.

B. Patient Recovery Classification

The objective of this ML task is to classify whether a patient has recovered from COVID-19 symptoms based on their device

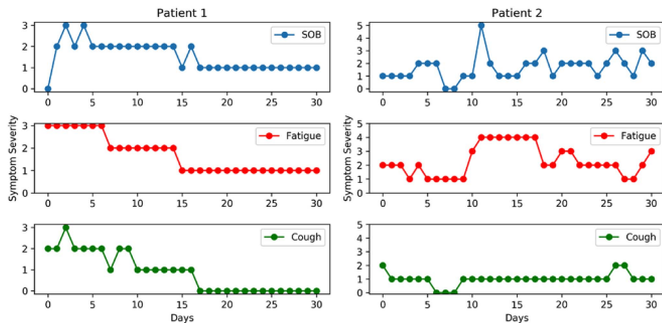


Fig. 3. Symptom severity progression for two COVID-19 patients. Patient 2's symptom severities decrease by day 7 and then sharply increase again after day 10. The shortness of breath (SOB), fatigue, and cough severities correspond to questions 3-5 of the symptom tracker.

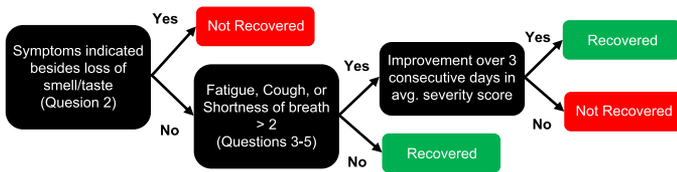


Fig. 4. Labeling logic for patient recovery classification based on symptom tracker questionnaire responses.

data. This binary classification model can provide healthcare workers with automated insights into the recovery status of their infected patients and bypass the need for manual daily symptom tracking which relies on patient compliance. To the best of our knowledge, there is no clear definition for full recovery from COVID-19. The US CDC recommends removal of isolation for COVID-19 infection when a patient's symptoms have significantly improved, they have been afebrile for at least 24 hours in the absence of fever-reducing medications, and it has been at least 10 days since symptom onset [23]. However, it is now understood some patients can suffer from ongoing symptoms from COVID-19 for weeks and even months [24]. Unlike symptom severity which can be identified by patients themselves, recovery is a gradual, subtle, and implicit process. In this task, we classify whether a patient has recovered from the COVID-19 symptoms collected by our symptom tracker app. Most patients experienced a steady decline in symptom severities, however, some patients initially appeared to recover and then had symptoms re-appear. Fig. 3 displays the symptom severity progression for the first 30 days for two different COVID-19 patients in terms of shortness of breath (SOB), fatigue, and cough. Patient 1 is an example of a patient who experienced a steady recovery. Patient 2, however, demonstrates a complicated symptom progression. The symptom severities for this patient declined by day 7 and then sharply worsened after day 10, especially for SOB and fatigue. All three symptoms linger for this patient for over a month.

A binary label is generated on a daily basis for each patient: recovered (0) or not recovered (1). The labelling logic for patient recovery was developed in collaboration with UCSD Health doctors and is displayed in Fig. 4. If symptoms are present besides loss of taste/smell (Question 2), label as not recovered (1). We do not consider loss of smell/taste because these symptoms have been shown to linger after a patient has recovered from COVID-19 [25]. If no symptoms are marked for Question 2

TABLE III  
STATISTICS FOR LABEL COUNT PER PATIENT

	Mean	Std.	Max	Min	Median
Not Recovered	24	29	85	0	16
Recovered	21	26	76	0	16

and fatigue/cough/shortness of breath severity is  $\leq 2$  (Questions 3-5), label as recovered (0). If fatigue/cough/shortness of breath severity is  $> 2$  but there is an improvement over 3 consecutive days in severity scores, label as recovered (0). In order to accommodate for complex cases such as Patient 2 in Fig. 3, in which there may be a day labeled as recovered (0) between days labeled as not recovered (1), we apply the following logic. If a patient is labeled as recovered (0) for 7 consecutive days, all the following labels are also marked as recovered (0). Otherwise, the recovered (0) days shorter than 7 days are reverted to non-recovered (1) days. This ensures there are no "recovered" days between "not recovered" days and vice versa. The statistics of the symptom tracker labels are shown in Table III. The average number of "not recovered" and "recovered" samples per patient is 24 and 21, respectively. The median number of "not recovered" and "recovered" samples per patient is 16 for both. This difference in mean and median is the result of outlier patients who have a high amount of one label. There are 10 patients for which 90% of their labels are either "not recovered" or "recovered". Patients with few "not recovered" labels may be a result of being asymptomatic or a delay in joining the study after being infected and testing positive. Patients with few "recovered" labels remained symptomatic for the study duration. These labels are used for the patient recovery classification task. Note that the recovery classification technique proposed here can be used with any other labeling logic developed by other health care providers.

### C. Device Data and Preprocessing

The Garmin vivosmart4 includes a heart rate monitor, accelerometer, ambient light sensor, and blood oxygen saturation (SpO<sub>2</sub>) monitor. The device uses these sensors in order to calculate various health parameters, including lifestyle and vitals information. The device data is presented in Table IV. The Garmin API documentation provides a description of these parameters [22]. Lifestyle features include activity (steps, distance, floors, active time, etc.), stress (average stress, max stress, stress duration, etc.), sleep timing (duration, bed time, up time), and sleep stages (deep, light, REM, awake). Stress-related features are derived based on heart rate variability [22]. The variable length of time in between each heartbeat is regulated by the body's autonomic nervous system. The less variability between beats equals higher stress levels, whereas the increase in variability indicates less stress. As mentioned in the introduction, the researchers in [12] found that COVID-19 affected the number of daily steps and time asleep for patients in their study. This result motivates us to include all lifestyle features when training our patient recovery classification model. In addition to lifestyle factors, the vivosmart4 measures vitals data including heart rate and SpO<sub>2</sub>. The device is capable of manual SpO<sub>2</sub> spot checks during the day and 4 hours of continuous measurement during sleep. Since the symptoms data and patient recovery classification labels are generated daily, we aggregate the device

**TABLE IV**  
LIST OF GARMIN DEVICE FEATURES THAT OUR APPROACH USES

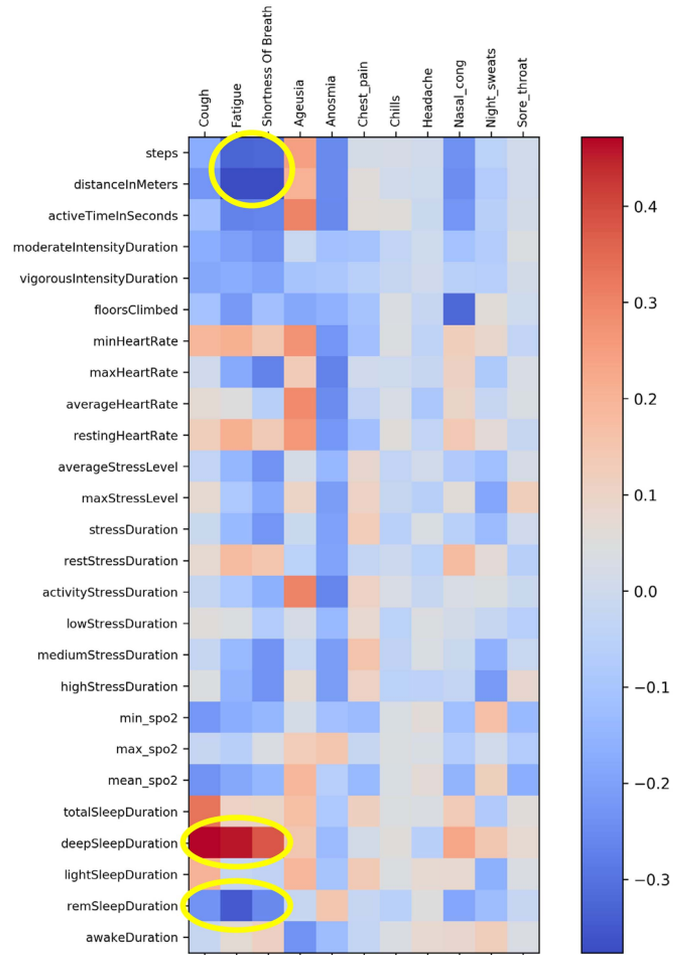
Features
Steps <sup>^</sup> , Distance, ActiveTime, ModerateIntensityDuration, VigorousIntensityDuration, FloorsClimbed, AverageStressLevel, MaxStressLevel, StressDuration, RestStressDuration, ActivityStressDuration, LowStressDuration, MediumStressDuration, HighStressDuration, SleepDuration <sup>^</sup> , BedTime* <sup>^</sup> , UpTime* <sup>^</sup> , DeepSleepDuration <sup>^</sup> , LightSleepDuration <sup>^</sup> , REMSleepDuration <sup>^</sup> , AwakeDuration <sup>^</sup> , MinHeartRate <sup>^</sup> , MaxHeartRate <sup>^</sup> , MeanHeartRate <sup>^</sup> , RestingHeartRate <sup>^</sup> , MinSpO <sub>2</sub> * <sup>^</sup> , MaxSpO <sub>2</sub> * <sup>^</sup> , MeanSpO <sub>2</sub> * <sup>^</sup>
Features marked with * require additional processing after receiving the data from garmin. Features marked with ^ are available in the dataset from [12] which we discuss in section IV-B.

**TABLE V**  
TOP 10 CORRELATIONS BETWEEN SYMPTOMS AND DEVICE FEATURES

Symptom	Device Feature	Spearman Correlation
Cough	DeepSleepDuration	0.47
Fatigue	DeepSleepDuration	0.46
SOB	DeepSleepDuration	0.38
SOB	DistanceInMeters	-0.38
Fatigue	DistanceInMeters	-0.37
Fatigue	REMSleepDuration	-0.34
Cough	TotalSleepDuration	0.33
Fatigue	Steps	-0.33
SOB	Steps	-0.32
Nasal Congestion	FloorsClimbed	-0.32

data features for each day. The Garmin Health API provides summarized activity, sleep, stress, and heart rate features daily. The features in Table IV marked with a \* require additional processing after receiving the data from Garmin. These include BedTime, UpTime, MaxSpO<sub>2</sub>, MinSpO<sub>2</sub>, and MeanSpO<sub>2</sub>. The BedTime and UpTime features are encoded as the number of seconds before or after midnight (e.g., 11:30 PM bed time is encoded as -1800 seconds, 8:00 AM wake time is encoded as 28800 seconds). Since only the continuous SpO<sub>2</sub> data is available through the Garmin API, we transform the SpO<sub>2</sub> time series each day into the MaxSpO<sub>2</sub>, MinSpO<sub>2</sub>, and MeanSpO<sub>2</sub> features displayed in Table IV. Note that a subset of the features is marked with ^ in Table IV indicating they are available in the dataset from [12] which we discuss in Section IV-B. Once the device data is aggregated for each day, we match it with the corresponding patient recovery label to form patient-day samples. Each patient-day sample consists of the recovery label and the summarized lifestyle and vitals features for one patient’s day in the study. Note that symptoms data are not directly used as part of the training data, but rather to generate the daily patient recovery labels.

Fig. 5 displays a heatmap of the correlation between the aggregated daily lifestyle/vitals features and symptoms data for our study cohort. We use Spearman correlation because the symptom variables are not continuous. Spearman evaluates the monotonic relationship between two continuous or ordinal variables [26]. The color of each heatmap square describes the magnitude and directionality of the correlation. Darker red squares correspond to a stronger positive correlation while darker blue squares correspond to a stronger negative correlation. Table V displays



**Fig. 5.** Spearman correlation between lifestyle/vitals and symptoms. Notable correlations are circled in yellow.

the top 10 most significant correlations between symptoms and device features and in Fig. 5 we circle notable correlations in yellow. These include distance and steps vs. fatigue and shortness of breath (SOB) severity, and deep and REM sleep vs. cough and fatigue severity. The correlations for distance vs. SOB and fatigue are  $-0.38$  and  $-0.37$ , respectively. The correlations for steps vs. SOB and fatigue are  $-0.32$  and  $-0.33$ , respectively. It is sensible that distance and steps are negatively correlated with cough and SOB severity. A patient is less likely to be active if their symptom severities are higher. Deep and REM sleep duration are positively and negatively correlated, respectively, with cough, fatigue and SOB severity. The most significant correlation is deep sleep vs. cough, which has a

correlation of 0.47. REM sleep is most correlated with fatigue, with a correlation of  $-0.34$ . According the American Academy of Sleep Medicine, as the immune system fights infection, the amount of time spent in REM sleep is decreased while deep sleep is increased [27]. This is because it is during deep sleep that many reparative bodily processes occur. This validates the directionality of the correlations between REM/deep sleep and symptom severities. While the individual correlations between other lifestyle/vitals features and symptoms are not as prominent, the heatmap in Fig. 5 indicates that a combination of these features can provide useful information about symptom severity when training the ML model. Overall, these correlation observations help motivate our ML approach to patient recovery classification based on device data.

#### D. Random Forest and Personalization

We train multiple ML classifiers in order to determine which is most effective at modelling the patient recovery task, as described in Section IV-A. As indicated in Table VI, the Random Forest (RF) model results in the best performance during LOSO CV. In this section, we discuss the operation of the RF model and our personalization technique.

RF is an ensemble model that aggregates a collection of decision trees in order to reduce overfitting and the resulting high variance in prediction [28]. To do this, RF utilizes bootstrap aggregation (bagging) and feature bagging. RF produces bootstrap datasets that are randomly and independently drawn with replacement from the training dataset. Each bootstrap dataset has the same size as the original training set and is used to train a decision tree. Bootstrap aggregation in RF averages the prediction of all decision trees which greatly reduces the variance compared to a single decision tree. Moreover, since individual trees generated in the bagging process are identically distributed, the expected prediction of RF is the same as the expected prediction of individual trees. Combining the above facts, RF has a lower variance than individual trees, while its bias remains the same [29]. RF further reduces the correlation between its member decision trees by introducing feature bagging, which randomly selects a subset of features when constructing each tree. In addition, RF is known to perform well even when using redundant or irrelevant features. Since we utilize multiple lifestyle and vitals features for model training, it is possible that some features do not provide useful information. Since RF is more robust to noisy features as compared to the other models [30], redundant or irrelevant features will not greatly impact performance.

Multiple studies that focus on ML for health applications have shown that model personalization is a key step in improving performance due to the physiological differences between patients [31], [32], [33], [34]. In this study, we observe that vitals and lifestyle factors vary among patients and propose a RF-based personalization technique to tune the model to each patient. Our technique involves including the first  $k$  days of labeled data from the test patient in the training set. In the traditional RF bootstrapping process, each training sample has uniform weight, which means each data sample is resampled with the same probability. To emphasize the test patient's calibration samples during model training, we assign a greater weight to these  $k$  samples using the Weighted Bootstrapping algorithm [35]. In order to implement this algorithm, a vector of sample weights  $\mathbf{W} = w_1, w_2, \dots, w_N$  is maintained where  $N$  is

the total number of training samples. Weights  $w_1, \dots, w_k$  correspond to the  $k$  personalization samples from the test patient and are given larger values. Weights  $w_{k+1}, \dots, w_N$  correspond to the data samples from the remaining patients used for training and are assigned lower values. The operation of the Weighted Bootstrapping algorithm is as follows [35]: First, a new bootstrap dataset for one decision tree is initialized. Then, the weights in  $\mathbf{W}$  are mapped into the interval  $[0, \sum_{j=1}^N w_j]$  with subintervals  $I_1, I_2, \dots, I_N$ . The length of each subinterval is proportional to the value of its weight. Next, each data sample is drawn using subintervals  $I_1, I_2, \dots, I_N$  and the uniform distribution function. The process repeats  $N$  times such that the size of all bootstrap datasets equals that of the original dataset. Consequently, the samples with higher weights are more likely to appear in each bootstrap dataset. In Section IV-B, we compare the performance for different values of  $k$  and different values of  $w_1, \dots, w_k$ . Fig. 6 displays a block diagram of our proposed RF personalization technique. After preprocessing each patient's data, Hybrid-CV is carried out in which the training and test sets are split on a per patient basis and the first  $k$  days of test patient data are added to the training set as personalization samples, as shown in Fig. 6. These  $k$  samples are assigned greater weights, which are bolded in the figure, during weighted bootstrapping. After training, the model is evaluated on the remaining, future data samples of the test patient.

## IV. RESULTS AND DISCUSSION

In this section, we describe the experiment settings and present patient recovery classification results. We discuss the effects of our RF model personalization technique on performance and carry out feature analysis using Shapley Values in order to interpret what the model has learned. Finally, we provide a discussion on the challenges encountered during this study.

### A. Experiment Setting

We implement and evaluate our machine learning models using the Scikit-learn library in the python environment on an Intel i5 3.2GHz quad-core and 16GB RAM computer. Accuracy, sensitivity, specificity, and F1-score are calculated and used as our evaluation metrics for the patient recovery classification task. For this task, a negative and positive sample correspond to a "recovered" and "not recovered" patient-day sample, respectively. Accuracy returns an overall measure of how much the model is correctly predicting on the entire set of test data. Sensitivity and specificity measure the true positive and true negative rate, respectively. F1 score is calculated as the harmonic mean of precision and recall (sensitivity) and is used to find the best trade-off between the two quantities [36]. As a result, we use F1 score for deciding the top performing model.

We carry out LOSO CV to mirror the clinically relevant use-case scenario of diagnosis for newly infected subjects [43]. LOSO CV separates the data into train and test sets on a per patient basis in order to simulate the practical application. This data split ensures that data from the same patient does not appear in both the training and testing sets. We use LOSO CV to compare the performance of different ML models. We then carry out Hybrid-CV, in which a specified number of samples from the test patient are included in the training set. These personalization samples are not included in the test set to ensure there is no overlap between train and test sets at the

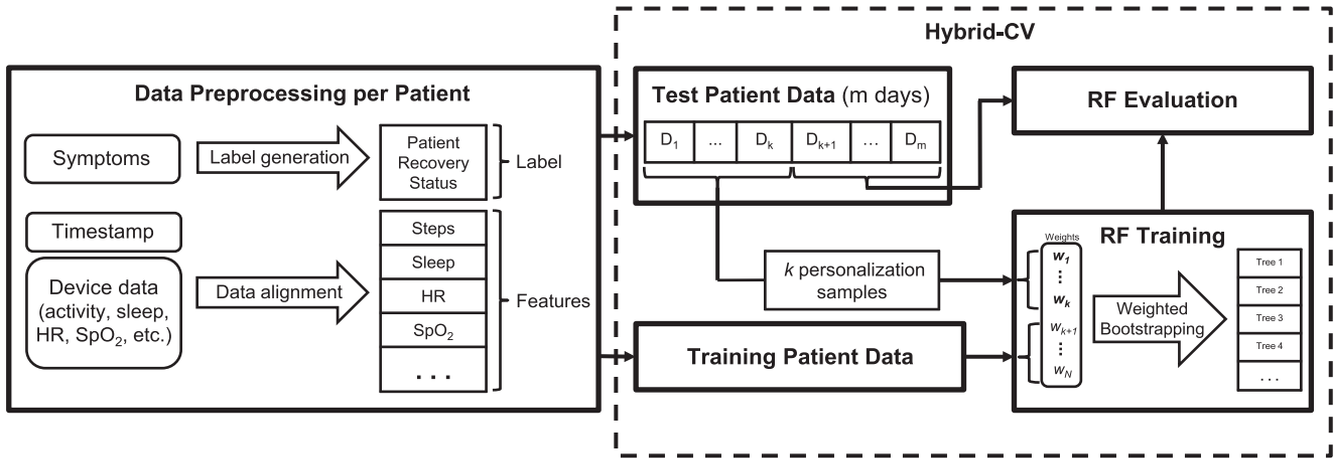


Fig. 6. Block diagram of our proposed RF personalization approach. After data preprocessing, the first  $k$  samples from the test patient are included in the training set during Hybrid-CV. These samples are assigned larger weights, which are bolded in the figure, during weighted bootstrap aggregation. After training, the model is evaluated on the remaining test patient data samples.

TABLE VI  
COMPARISON OF ML MODEL PERFORMANCE FOR LOSO CV

Model	Acc	Sens	Spec	F1
LR	0.60	0.61	0.52	0.61
ANN	0.59	0.62	0.62	0.63
SVM	0.54	0.61	0.59	0.62
KNN	0.55	0.51	0.68	0.60
LSTM-16	0.63	0.56	0.71	0.61
LSTM-32	<b>0.64</b>	<b>0.64</b>	0.60	0.64
RF	0.59	0.52	<b>0.78</b>	<b>0.66</b>

sample level. We compare how performance is affected by applying varying levels of personalization using our RF-based personalization technique described in Section III-D. Since the number of samples for each patient is different based on their participation in the study, the training and testing sets will vary in size for both CV experiments. Instead of averaging the results over each data split, we save the model predictions for each data split and calculate metrics over all predictions. This ensures that each patient-day contributes equal weight to the final result.

In the LOSO CV experiment, we compare RF with the following ML models: logistic regression (LR) [37],  $k$ -nearest neighbors (KNN) [38], support vector machine (SVM) [39], artificial neural network (ANN) [40], and long short-term memory (LSTM) neural network [41]. Model hyperparameter tuning is performed with each training set using a randomized search over a predefined hyperparameter grid for each model. Since LSTM models take sequential data as input, we organize the lifestyle and vitals features into sequential data samples using a window length of 7 days and a step size of 1 day. A step size of 1 day is used to extract the maximum number of samples. As a result, each input sample has a dimension of  $(7, N_{features})$  where  $N_{features}$  represents the number of lifestyle and vitals features. The patient recovery label for the last day of each window is assigned to each input sample. We train the LSTM as a

many-to-one model, as opposed to a many-to-many model, since the application of this method is only concerned with estimating whether the patient is recovered or not for the current day. In addition, training the LSTM to estimate one label at a time matches the process for the other ML models, resulting in a fairer comparison. We carry out two LSTM experiments using 16 and 32 hidden units for the LSTM layer followed by a fully connected layer with 1 output unit. For these experiments, we train the models using the Adam optimizer [42] and a dropout rate of 50% to reduce overfitting. For the LSTM layers, we use a sigmoid activation function for the input, forget and output gates, and a hyperbolic tangent (tanh) activation function for the cell state and hidden state. The fully connected layers use a sigmoid activation function and we use binary cross entropy loss as the loss function. We experimented with different numbers of training epochs and batch sizes and found that 25 epochs and a batch size of 32 resulted in the best performance.

### B. Patient Recovery Classification Results

Accuracy, sensitivity, specificity, and F1-score for each ML model during LOSO CV are presented in Table VI. The LSTM-32 model achieves the highest accuracy and sensitivity, both equal to 0.64, while the RF model achieves the highest specificity and F1 score equal to 0.78 and 0.66, respectively. As described in the experiment setting, we use F1 score for deciding the top performing model since this metric calculates the tradeoff between precision and sensitivity. Since RF achieves the highest F1 score, we conclude that RF is the best performing model for patient recovery classification. We attribute the RF's top performance to its ability to reduce the variance in prediction via the bagging process and its robustness to redundant or irrelevant features. The LSTM-32 model is the second-best performing model, indicating that meaningful temporal information exists in the data for estimating recovery from COVID-19. Since RF is the top performer, we use this model in the next experiment to understand how the number of personalization samples impacts RF performance.

Next, we discuss the results of the Hybrid-CV experiment. As mentioned in the experiment settings, LOSO CV separates the data into train and test sets on a per patient basis. Since

TABLE VII

HYBRID-CV RESULTS USING DIFFERENT LEVELS OF PERSONALIZATION

Personalization Samples	Acc	Sens	Spec	F1
0	0.59	0.52	0.78	0.66
1	0.63	0.59	0.75	0.70
2	0.67	0.66	0.71	0.75
3	0.72	0.73	0.68	0.79
4	0.80	0.86	0.64	0.86
5	0.82	0.89	0.63	0.88

physiology and lifestyle differ between patients, we apply varying levels of personalization during the Hybrid-CV experiment. We implement our RF-based personalization technique by including the first 1-5 days of test patient data in the training set. These personalization samples are assigned a larger weight so that they are sampled more frequently during the bootstrap aggregation step. Table VII displays the results for different amounts of personalization. Evidently, the classification results are worse when no personalization is applied. The accuracy, sensitivity, specificity, and F1-score are 0.59, 0.52, 0.78, and 0.66, respectively, when no personalization is applied. As personalization samples are included in the training set, accuracy, sensitivity, and F1-score increase, while specificity decreases. When using 5 personalization samples, the accuracy, sensitivity, specificity, and F1-score are 0.82, 0.89, 0.63, and 0.88, respectively. Since the personalization samples for each patient correspond to their first 1-5 days in the study, these samples are primarily labeled 1 or “not recovered”. This means that as more personalization samples are included in the training set, the model can increasingly learn the infected baseline of the patient based on their vitals and lifestyle data. This causes the sensitivity to increase since the model will be able to increasingly correctly classify a patient who has not recovered. This corresponds to increasing true positives (classifying a patient as not recovered when they are indeed not recovered) while minimizing false negatives (classifying a patient as recovered when they are not recovered). As the sensitivity increases, the specificity decreases. Since the model is increasingly tuned to classify a patient as not recovered, this will result in more false positives and a lower specificity. For this ML task, false positives are more acceptable than false negatives. Classifying a patient as not recovered when they are recovered is less harmful than classifying a patient as recovered when they are not recovered. Overall, adding personalization samples increases the model performance. When applying this personalization technique to a new patient, the first few days will involve data collection without any classifications from the ML model. After this initial data collection, the personalized model will provide estimations with improved accuracy, sensitivity, and F1-score. The results demonstrate the potential for ML-assisted remote patient monitoring to supplement traditional manual monitoring tools, like daily manual symptom tracking.

The results presented in Table VII are generated by setting the bootstrap aggregation weights for the personalization samples to 10. This means these samples are 10 times more likely to be sampled during the RF weighted bagging process. In Table VIII,

TABLE VIII

PERFORMANCE COMPARISON WHEN APPLYING DIFFERENT RF BOOTSTRAP AGGREGATION WEIGHTS TO 5 PERSONALIZATION SAMPLES

Bootstrap Aggregation Weights	Acc	Sens	Spec	F1
1	0.70	0.69	0.73	0.77
10	0.82	0.89	0.63	0.88
100	0.81	0.88	0.62	0.87

TABLE IX

EVALUATION OF PROPOSED METHOD ON OPEN DATASET FROM [12]

	Acc	Sens	Spec	F1
W/O Personalization	0.49	0.33	0.73	0.44
W/ Personalization (5 samples)	<b>0.61</b>	<b>0.55</b>	<b>0.67</b>	<b>0.61</b>

we compare how classification performance is affected by applying different bagging weights to 5 personalization samples. We set the weights to 1, 10 and 100. Using a bagging weight of 1 means the personalization samples have the same probability of being sampled as the training data from other patients. Evidently, a bagging weight of 1 produces worse performance with an accuracy, sensitivity, specificity, and F1-score of 0.7, 0.69, 0.73, and 0.77, respectively. In this case, the personalization samples are not emphasized and the model is not effectively calibrated. Increasing the bagging weight from 10 to 100 does not improve model performance. This indicates that at a certain weight, the personalization samples are sampled frequently enough during bagging to effectively calibrate the model. Further increasing the bagging weight does not provide additional utility in model personalization.

In order to extend the evaluation of our proposed method, we applied our approach to the dataset collected in [12]. This dataset includes sleep, heart rate and steps data collected from a wearable device, and the date of first symptoms and date of recovery which are manually recorded by each patient. Since this dataset does not include SpO<sub>2</sub>, stress or activity (besides steps) data, the number of features is significantly less than our own dataset (12 vs. 28). In Table IV, features marked with ^ are available in the dataset in [12]. We labelled all days between the start of symptoms and recovery dates as “not recovered” and all days after the recovery date as “recovered.” We then combined these labels with the corresponding device features to create the dataset in the same manner as our experiment setting. After these data processing steps, 15 patients had sufficient data to be included in this experiment. Table IX displays the results when applying our method to this dataset. We train a Random Forest model with and without personalization and calculate the accuracy, sensitivity, specificity, and F1-score. We use 5 samples when applying our personalization technique and observe that the performance significantly improves compared to the non-personalized results. With personalization, our approach achieves an accuracy, sensitivity, specificity, and F1-score of 0.61, 0.55, 0.67, and 0.61, respectively. Evidently, the performance metrics are not as good for this dataset. This may be due to the limited feature set and inaccurate recovery dates

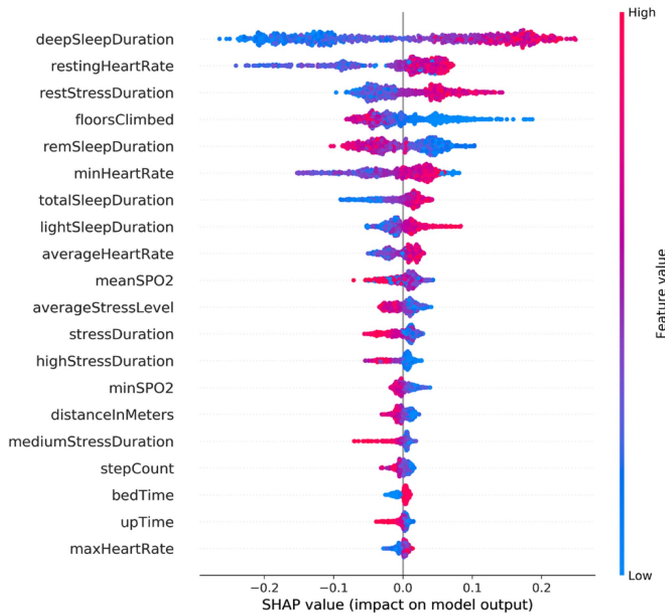


Fig. 7. Summary of Shapley top features where each point corresponds to a data sample. The x-axis represents a feature's impact on model output. Positive SHAP values push the model to output 1 or "not recovered".

recorded by patients. We observe similar patterns in the results compared with our own dataset which include that there is a performance enhancement when applying our personalization technique. Overall, these consistent observations between our dataset and the dataset in [12] indicate that our proposed approach is not only applicable to our dataset, but can potentially be applied to different datasets collected in clinical practice.

### C. Model Interpretability Via Shapley Value Analysis

Next, we utilize Shapley Values [44], [45] in order to determine which lifestyle and vitals features have the most significant effect on model classification for our dataset. Shapley Value analysis is a model-agnostic interpretation method derived from game theory. Given a set of feature values and a trained machine learning model, the estimated Shapley value indicates how each feature contributes to the model's classification. We use the tree SHAP (SHapley Additive exPlanations) framework [46], [47], which is optimized for tree-based models, to interpret the output of the RF model for patient recovery classification. Fig. 7 displays the Shapley results where the features are ranked from the top to bottom based on their impact on the model's output. Each point on the plot corresponds to an individual data sample and represents the contribution from the feature listed on the Y-axis to the RF's classification. The placement on the X-axis represents the amount of positive/negative contribution to the classification. Positive contribution corresponds to pushing the model to estimate that a patient is not recovered. The color of each point represents the actual value of the feature (red is high while blue is low). The top two features based on Shapley analysis include deep sleep duration and resting heart rate. Higher values of deep sleep duration (colored in red) contribute to a positive, or not recovered, classification. This observation aligns with the correlation analysis presented in Section III-C. As mentioned earlier, deep sleep increases when a patient is sick since

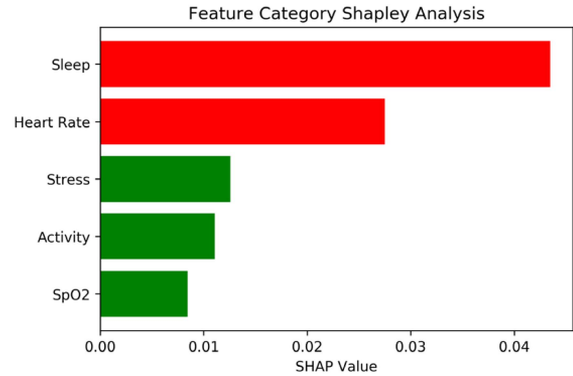


Fig. 8. Impact of feature categories on model output. Features are grouped into 5 categories and a categorical SHAP score is calculated. Red or green bars indicate that an increase in the category's feature values pushed the model to output "not recovered" or "recovered," respectively.

this is when many reparative bodily processes occur. Increased resting heart rate also contributes to a positive classification by the RF model. This relationship makes sense since resting heart rate will decrease as a patient recovers. Additional observations include that a lower number of floors climbed contributes to a positive classification while an increased mean SpO<sub>2</sub> contributes to a negative, or recovered, classification. Both relationships are sensible, as a patient who has not recovered will be less active and a patient who has recovered will have a higher SpO<sub>2</sub>.

In addition to analyzing the impact of individual features, we grouped the features into 5 categories (Activity, Sleep, Stress, Heart Rate and SpO<sub>2</sub>) and investigated their impact on model output. A SHAP score for each category was calculated as the average of the absolute SHAP values for the features in that category. Fig. 8 displays the ranking of feature categories based on their categorical SHAP score. We also examined whether, on average, an increase in the feature values for each category pushed the model to estimate "recovered" or "not recovered." In Fig. 8, a red colored bar indicates that an increase in the category's feature values pushed the model to output "not recovered." A green colored bar indicates that an increase in the category's feature values pushed the model to output "recovered." Evidently, the sleep category had the most significant impact on model output. An increase in feature values in the sleep and heart rate categories pushed the model to estimate "not recovered" (red bars) while an increase in feature values in the stress, activity and SpO<sub>2</sub> categories pushed the model to estimate "recovered" (green bars). Overall, the individual feature and feature category Shapley analysis demonstrates that our model can learn clinically relevant relationships between device data and the status of patients. The interpretability of a ML model is necessary for humans to understand what the model has learned, especially in medical applications.

### D. Limitations and Research Challenges Encountered

In this section, we discuss limitations to our proposed approach and challenges faced while implementing this study. One limitation in our approach is that patients were only enrolled and provided devices for data collection after testing positive for COVID-19. It is likely that some patients started experiencing symptoms before going for a COVID-19 test. This meant we

were not able to collect symptoms and wearable data during the initial days of the infection. In order to ensure that data can be collected before and during the onset of COVID-19 infection, participation could be made available to a larger number of patients that already own a wearable device. After testing positive for COVID-19, a patient could immediately enroll and begin sharing both past and current data. Another limitation to our approach is that the RF model does not process data sequentially while the progress of COVID-19 is sequential. In this work, we experimented with LSTM, a popular temporal model, however, found its performance to be worse than RF. Training an LSTM requires significantly more data since neural networks are highly prone to overfitting when the underlying dataset size is small [48], [49]. In order to fully utilize temporal relationships in the data, we plan to further investigate sequence modeling with additional data in our future work. This will include implementing many-to-many sequence models using different time windows to learn temporal progression along with the label. In addition, a larger dataset can enable the use of additional features such as patient demographic information. The model may learn relationships between COVID-19 recovery and demographic data such as age, gender, and ethnicity.

Concern over privacy was an issue encountered during recruitment for this study. As mentioned in Section III-A, we recruited patients from both the UCSD Health and Neighborhood Healthcare (NH) COVID-19 telemedicine clinics. NH is a community clinic that primarily provides care to underserved populations. In order to increase accessibility to our study, we developed a Spanish version of our symptom tracker app with assistance from NH. Overall, we experienced more difficulty recruiting from NH. One reoccurring reason why NH patients did not want to partake in our study included a concern over privacy. Certain patients expressed discomfort over wearing the device 24/7 due to concerns of being tracked. Our recruitment personnel would highlight that the device does not collect any location data, however, certain patients still declined participation. The above challenge encountered during our study showed that privacy concerns and lack of trust in wearables may further limit access and use of digital technologies by underserved populations, contributing to an increased digital divide in healthcare. As healthcare begins to rely more on digital technologies, these concerns must be addressed in order to ensure equal access to high quality healthcare [50].

## V. CONCLUSION

In this paper, we propose an intelligent remote monitoring platform, namely eCOVID, for enhanced COVID-19 ambulatory care. Based on data collected from our study with the UCSD Health and Neighborhood Healthcare COVID-19 telemedicine clinics in San Diego County, we demonstrate correlations between automatically collected wearable data and manually entered symptom data. We propose a novel ML approach to estimate whether a patient has recovered from COVID-19 symptoms based on the automatically collected wearable data. Our results demonstrate that ML-assisted remote monitoring using wearable data can supplement or be used in place of manual daily symptom tracking which relies on patient compliance.

By developing and demonstrating the ability to track patient recovery status remotely, our approach can enable more optimal care of COVID-19 ambulatory patients at scale. Care teams will be able to track patient recovery efficiently through

automatically generated and updated dashboards instead of the current practice of manual symptom tracking and phone calls, the latter becoming ineffective when there is a surge in cases. This shift can lead to significant improvement in the efficiency and scalability of ambulatory patient care, while at the same time enabling savings in human and equipment resources. Moreover, the approach can be used for providing scalable and efficient care for future pandemic and epidemic challenges.

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*Data Availability:* The de-identified data used in this study can be downloaded from the study data repository on IEEE DataPort (<https://iee-dataport.org/>). This includes the symptom tracker responses and data collected from Garmin devices.

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