Wearable Finger Pulse Oximetry for Continuous Oxygen Saturation Measurements During Daily Home Routines of Patients With Chronic Obstructive Pulmonary Disease (COPD) Over One Week: Observational Study

Joren Buekers1,2, MSc; Jan Theunis1, MSc; Patrick De Boever1,3, PhD; Anouk W Vaes4, PhD; Maud Koopman4, MSc; Eefje VM Janssen4, MSc; Emiel FM Wouters4,5, PhD; Martijn A Spruit4,5,6,7, PhD; Jean-Marie Aerts2, PhD

1Health Unit, Flemish Institute for Technological Research (VITO), Mol, Belgium
2Measure, Model & Manage Bioresponses, Department of Biosystems, KU Leuven, Leuven, Belgium
3Centre for Environmental Sciences, Hasselt University, Diepenbeek, Belgium
4Department of Research and Education, Centre of Expertise for Chronic Organ Failure (CIRO), Horn, Netherlands
5Department of Respiratory Medicine, Maastricht University Medical Centre, Maastricht, Netherlands
6Rehabilitation Research Center (REVAL), Biomedical Research Institute (BIOMED), Faculty of Rehabilitation Sciences, Hasselt University, Diepenbeek, Belgium
7School of Nutrition and Translational Research in Metabolism (NUTRIM), Maastricht University Medical Centre, Maastricht, Netherlands

Corresponding Author:
Jean-Marie Aerts, PhD
Measure, Model & Manage Bioresponses
Department of Biosystems
KU Leuven
Kasteelpark Arenberg 30
Leuven, 3000
Belgium
Phone: 32 16321434
Email: jeanmarie.aerts@kuleuven.be

Abstract

Background: Chronic obstructive pulmonary disease (COPD) patients can suffer from low blood oxygen concentrations. Peripheral blood oxygen saturation (SpO2), as assessed by pulse oximetry, is commonly measured during the day using a spot check, or continuously during one or two nights to estimate nocturnal desaturation. Sampling at this frequency may overlook natural fluctuations in SpO2.

Objective: This study used wearable finger pulse oximeters to continuously measure SpO2 during daily home routines of COPD patients and assess natural SpO2 fluctuations.

Methods: A total of 20 COPD patients wore a WristOx2 pulse oximeter for 1 week to collect continuous SpO2 measurements. A SenseWear Armband simultaneously collected actigraphy measurements to provide contextual information. SpO2 time series were preprocessed and data quality was assessed afterward. Mean SpO2, SpO2 SD, and cumulative time spent with SpO2 below 90% (CT90) were calculated for every (1) day, (2) day in rest, and (3) night to assess SpO2 fluctuations.

Results: A high percentage of valid SpO2 data (daytime: 93.27%; nocturnal: 99.31%) could be obtained during a 7-day monitoring period, except during moderate-to-vigorous physical activity (MVPA) (67.86%). Mean nocturnal SpO2 (89.9%, SD 3.4) was lower than mean daytime SpO2 in rest (92.1%, SD 2.9; P<.001). On average, SpO2 in rest ranged over 10.8% (SD 4.4) within one day. Highly varying CT90 values between different nights led to 50% (10/20) of the included patients changing categories between desaturator and nondesaturator over the course of 1 week.

Conclusions: Continuous SpO2 measurements with wearable finger pulse oximeters identified significant SpO2 fluctuations between and within multiple days and nights of patients with COPD. Continuous SpO2 measurements during daily home routines of patients with COPD generally had high amounts of valid data, except for motion artifacts during MVPA. The identified
fluctuations can have implications for telemonitoring applications that are based on daily SpO₂ spot checks. CT90 values can vary greatly from night to night in patients with a nocturnal mean SpO₂ around 90%, indicating that these patients cannot be consistently categorized as desaturators or nondesaturators. We recommend using wearable sensors for continuous SpO₂ measurements over longer time periods to determine the clinical relevance of the identified SpO₂ fluctuations.

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KEYWORDS COPD; oxygen saturation; finger pulse oximeter; wearable sensor; nocturnal desaturation; telemonitoring

Introduction

Chronic obstructive pulmonary disease (COPD) is a highly prevalent lung disease that is characterized by persistent airflow limitation due to a mixture of obstructive bronchiolitis and emphysema [1]. Morbidity and mortality of COPD are high and still increasing [2], leading COPD to become the third-leading cause of death worldwide by 2030 [3]. COPD patients can suffer from low blood oxygen concentrations due to gas exchange abnormalities [1]. Hypoxemia during the night (ie, nocturnal desaturation) is also common in patients with COPD [4-6] due to nocturnal alveolar hypoventilation and ventilation-perfusion mismatching [7]. Hypoxemia can worsen with increasing disease severity [8]. Furthermore, it has been shown that hypoxemia is associated with lower exercise tolerance; decreased quality of life; increased risk for exacerbations, defined as “a sustained worsening of the patient's condition, from the stable state and beyond normal day-to-day variations” [9]; and higher risks of death [8].

The presence of hypoxemia can be assessed by invasive blood gas analyses. A noninvasive method to assess peripheral blood oxygen saturation (SpO₂) is pulse oximetry. Spot check SpO₂ measurements with thresholds of 88%-92% have been suggested for the detection of hypoxemia [1,10,11]. In telemonitoring applications, daily SpO₂ spot checks are used to raise alerts for exacerbations when SpO₂ spot check values drop below predefined SpO₂ thresholds [12]. Nocturnal desaturation is usually defined as having an SpO₂ value below 90% for more than 30% of the time in bed, measured during one [6,13-16] or two nights [5,17-20].

The current SpO₂ monitoring strategies do not take into account natural fluctuations in SpO₂. Sampling SpO₂ with such a low time frequency or during such a short time period may thus lead to classification errors (ie, hypoxicemic or nonhypoxicemic and nocturnal desaturator or nondesaturator) or false alerts in telemonitoring applications [12]. Wearable finger pulse oximeters provide the possibility to collect SpO₂ data at higher time frequencies and over longer time periods. This makes it possible to assess and account for oxygen saturation fluctuations in patients with COPD. Therefore, the objective of this study was to use wearable finger pulse oximeters to continuously measure SpO₂ during daily home routines of COPD patients and assess natural SpO₂ fluctuations. We hypothesized that significant natural SpO₂ fluctuations are present within and between multiple days and nights, which may lead to classification errors (ie, nocturnal desaturator or nondesaturator) or false alerts in telemonitoring applications.

Methods

Study Design and Participants

Figure 1 provides a general overview of the methods applied in this single-center, 1-week observational study. COPD patients at Global Initiative for Chronic Obstructive Lung Disease stages II-IV (GOLD II-IV) were recruited at the Centre of Expertise for Chronic Organ Failure (CIRO), a COPD treatment center located in Horn, the Netherlands, during a standard baseline assessment prior to pulmonary rehabilitation [21]. The target sample size was set a priori to 20 patients. COPD patients were eligible to enroll in the study based on the following criteria: (1) clinically stable (ie, no exacerbation in the past 4 weeks), (2) no rollator use, and (3) no long-term oxygen therapy. Patients that were interested in participating were called a few days after the baseline assessment to schedule a home visit for the delivery of the wearable sensors. During this visit, the functioning of the sensors and data acquisition protocol were explained, written informed consent was obtained, and a new visit was planned for collection of the sensors at the end of the study period. During the 7-day study period, which took place before the start of pulmonary rehabilitation, two phone calls were made to resolve potential technical difficulties. Demographics, oxygen partial pressure in arterial blood (PaO₂), postbronchodilator pulmonary function data (ie, forced expiratory volume in 1 second [FEV₁], forced vital capacity [FVC], and transfer factor for carbon monoxide [TLCO]), 6-minute walking distance (6MWD), and COPD assessment test (CAT) results were collected during the standard baseline assessment at CIRO. The study was approved by the Medical Research Ethics Committees United (MEC-U) (study approval number: NL58079.100.16) in the Netherlands and executed between December 2016 and April 2018.
Figure 1. General overview of the applied methods. COPD: chronic obstructive pulmonary disease; SpO$_2$: peripheral blood oxygen saturation; CT90: cumulative time spent with SpO$_2$ below 90%.

### Wearable Sensors and Protocol

Continuous SpO$_2$ measurements were performed for 1 week using a wearable finger pulse oximeter: WristOx$_2$ 3150 (Nonin Medical). Nonin oximeters have frequently been used for home monitoring of patients with COPD [12] and the WristOx$_2$ 3150 model complies with the International Organization for Standardization (ISO) standards ISO 10993-1 and ISO 80601-2-61. The manufacturer reports an accuracy of ±2% for SpO$_2$ measurements [22]. Sampling frequency was fixed at 1 Hz. Only the cumulative measurement time was visible for the participants. SpO$_2$ values were not shown on the WristOx$_2$ display to prevent patients from changing their behavior when deviating SpO$_2$ values would occur. Every participant received three WristOx$_2$ devices to deal with the limited battery life (ie, 48 hours of continuous measurements with one WristOx$_2$). Participants were instructed to wear the WristOx$_2$ on the index finger of their nondominant hand every night and as much as possible during the day, depending on their daily routines and comfort of wearing the finger clip. Raw data were stored on the internal memory of the WristOx$_2$ and downloaded at the end of the week using nVISION software, version 6.4.0.10 (Nonin Medical).

Simultaneously, actigraphy was performed with the SenseWear Armband (BodyMedia) for obtaining contextual information about physical activity levels and when the participants were lying down and/or asleep. The SenseWear Armband is a multisensory triaxial accelerometer, combining accelerometry with measurements of heat flux, galvanic skin response, and skin temperature. Based on these measurements, the armband provides information about, for example, energy expenditure (EE), expressed as metabolic equivalent of task (MET), or steps taken, while also indicating when the wearer is lying down and sleeping, at a standard sampling time of 1 sample per minute. Measurements of MET were used for classifying physical activity levels, whereas the indications of lying down and sleeping were used to separate daytime measurements from nocturnal measurements, as further described below. The SenseWear Armband has been shown to be accurate for measurements of both physical activity [23,24], except when using a rollator [24], and sleep estimations [25]. Patients were asked to wear the armband on the left upper arm, except when there was contact with water (eg, when taking a shower). Battery life of one SenseWear Armband was sufficient for continuous 24-hour measurements with a 1-minute sampling time during the whole week. Data were stored on the internal memory and downloaded at the end of the week using the BodyMedia SenseWear 8.1 software (BodyMedia).

### Data Preprocessing

Figure 2 visualizes the preprocessing steps. The SenseWear Armband indications about lying down and sleep were used to determine the time of going to bed in the evening and the time of getting out of bed in the morning. These time stamps were used to divide SpO$_2$ data into nocturnal and daytime data (see Figure 2A). Only full-night nocturnal SpO$_2$ measurements were retained for further analyses and daytime SpO$_2$ measurements were retained if there was at least one hour of SpO$_2$ measurements during that day. One hour of daytime measurements was considered sufficient to examine whether significant fluctuations occurred in SpO$_2$ values during the day. Previous studies examining intraday fluctuations only examined measurements of one hour or less [26-29]. No days or nights had to be excluded due to lacking SenseWear Armband measurements. Days with at least one hour of measurements in both the afternoon and evening were used for the comparison between afternoon (ie, 13:00-18:00) and evening (ie, 18:00-going to bed) SpO$_2$ values. No comparison was performed between SpO$_2$ values in the morning (ie, before 13:00) and the afternoon and evening, because SpO$_2$ measurements were often not performed before 13:00 (see Multimedia Appendix 1).
The raw SpO₂ data contained error values (ie, the number 500 was provided in the raw data file) when measurements were considered invalid by the algorithms of the WristOx₂ manufacturer (ie, orange dashes in Figure 2A). However, close inspection of the time series indicated that invalid data (eg, sudden low values) surrounded these error values. A data-cleaning algorithm was developed to exclude these invalid SpO₂ values as follows. First, small blocks of data (ie, less than 20 samples) between error values were excluded. Second, bigger blocks of data (ie, between 20 and 100 samples) between error values were excluded, only when the mean SpO₂ value of this data block was deviating more than 6% from the mean SpO₂ value of the full day or night under consideration: both steps are shown in Figure 2B. Thorough visual and raw data examination of both valid and invalid data led to the choice of this 6% threshold for excluding invalid blocks of data (eg, Figure 2B: around 17:00-18:00). Third, remaining outliers were excluded by down-sampling the data (see Figure 2C). Autocorrelation analyses indicated that a sampling time of 20 seconds was appropriate (see Multimedia Appendix 2) and all data were thus down-sampled by taking the median SpO₂ value of each consecutive 20-second block. By taking the median value, the effect of outliers was excluded. As a last step, small gaps of SpO₂ data (ie, 3 or fewer missing samples) were filled using linear interpolation (see Figure 2C). After application of the data-cleaning algorithm, all data were visually checked to ensure valid SpO₂ values were retained, while invalid values were removed.

Data Analyses
The SenseWear Armband indications about lying down and sleep were used to calculate the total night sleeping time (TNST) (ie, sum of all minutes indicated as sleep), wake time after sleep.
onset (WASO) (ie, sum of all minutes spent awake during the
time in bed, after the first onset of sleep), and sleep efficiency
(Seff) (ie, the ratio between TNST and time in bed) for every
separate night. Weekly averages of TNST, WASO, and Seff
were calculated to describe the sleep quality of the included
patients.

Daytime SpO\textsubscript{2} data were divided into daytime data during rest
(EE\textless 1.5 MET, while the patient was still awake), during
low-intensity physical activities (LIPAs) (1.5 MET<EE\textless 3
MET), and during moderate-to-vigorous intensity physical
activities (MVPAs) (EE>3 MET). Data quality of the continuous
SpO\textsubscript{2} measurements was assessed for both nocturnal and
daytime data based on the amount of valid data (ie, excluding
error values and cleaned values). Furthermore, the effect of
physical activity on the amount of valid data was examined by
comparing data quality during MVPA with data quality during
rest and LIPA (EE\textless 3 MET).

Mean SpO\textsubscript{2}, SpO\textsubscript{2} SD, and cumulative time spent with SpO\textsubscript{2}
below 90% (CT90) were calculated for every separate (1) day,
(2) day in rest, and (3) night. Hereafter, the weekly average and
weekly range (ie, the difference between the maximum and
minimum value over the different days or nights of the same
patient) of these features were calculated for every patient.

Intraday and intranight fluctuations were quantified as the
weekly average of the standard deviation of the SpO\textsubscript{2}
measurements. The range of SpO\textsubscript{2} values during the day in rest,
calculated as the difference between the maximum and minimum
SpO\textsubscript{2} values of that day in rest, was determined to indicate how
much spot-check values in telemonitoring applications could
differ depending on the moment of the measurement. Only
daytime SpO\textsubscript{2} in rest was considered because only at these
moments could spot checks have been performed in a
telemonitoring application. In addition, the difference between
mean SpO\textsubscript{2} values in rest in the afternoon (ie, between 13:00
and 18:00) and in the evening (ie, between 18:00 and going to
bed) was calculated for every day separately to examine
differences in SpO\textsubscript{2} baseline levels during the day.

Night-to-night and day-to-day SpO\textsubscript{2} fluctuations were quantified
as the weekly ranges of the three features (ie, mean SpO\textsubscript{2}, SpO\textsubscript{2}
SD, and CT90). Furthermore, we examined how many of the
included patients changed category between nocturnal desaturator and nondenaturator.

Statistical Analysis

Patient characteristics, weekly averages, and weekly ranges
were summarized for all patients as mean and SD. Paired-sample
t tests were used to test for differences between weekly averages
of nocturnal and daytime SpO\textsubscript{2} in rest. Pearson correlations
assessed the relationship between mean SpO\textsubscript{2} and patient
characteristics, intraday SpO\textsubscript{2} fluctuations, or intranight SpO\textsubscript{2}
fluctuations. A P value of <.05 was considered statistically
significant. All analyses were carried out in Jupyter Notebooks
(Project Jupyter) [30] using the Python 3.5 programming
language (Python Software Foundation) [31].

Results

Participants

A total of 21 out of 41 patients that were approached accepted
study participation. One patient suffered from an exacerbation
before the start of the 7-day study period and was excluded.
This finally led to the inclusion of 20 patients (14 males, 70%;
6 females, 30%) with moderate (8/20, 40%), severe (10/20,
50%), or very severe (2/20, 10%) COPD. General demographics,
postbronchodilator lung function, resting arterial blood gases,
6MWD, CAT score, and sleep-quality characteristics are
summarized in Table 1.
### Table 1. Characteristics of the 20 included patients with moderate-to-very severe COPD$^a$.

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>63 (8)</td>
</tr>
<tr>
<td>Body mass index (kg/m$^2$)</td>
<td>26 (4)</td>
</tr>
<tr>
<td>Forced expiratory volume in 1 second (L)</td>
<td>1.4 (0.5)</td>
</tr>
<tr>
<td>Forced expiratory volume in 1 second (% predicted)</td>
<td>48 (15)</td>
</tr>
<tr>
<td>Forced vital capacity (L)</td>
<td>3.8 (1.1)</td>
</tr>
<tr>
<td>Forced vital capacity (% predicted)</td>
<td>97 (20)</td>
</tr>
<tr>
<td>Transfer factor for carbon monoxide (% predicted)</td>
<td>50 (16)</td>
</tr>
<tr>
<td>Partial pressure of oxygen (kPa)</td>
<td>8.7 (1.7)</td>
</tr>
<tr>
<td>Partial pressure of carbon dioxide (kPa)</td>
<td>5.2 (0.6)</td>
</tr>
<tr>
<td>6-minute walking distance (m)</td>
<td>411 (59)</td>
</tr>
<tr>
<td>COPD assessment test score</td>
<td>21 (5)</td>
</tr>
<tr>
<td>Total night sleeping time (min)</td>
<td>411 (72)</td>
</tr>
<tr>
<td>Wake time after sleep onset (min)</td>
<td>74 (36)</td>
</tr>
<tr>
<td>Sleep efficiency (%)</td>
<td>83 (8)</td>
</tr>
</tbody>
</table>

$^a$COPD: chronic obstructive pulmonary disease.

### Continuous SpO$_2$ Measurements, Preprocessing, and Data Quality

An overview of the amount of SpO$_2$ measurements that were included for further analyses is provided in Table 2 and Multimedia Appendix 1. A total of 2 days and 2 nights of measurements from patient 4 were missing due to battery issues. A total of 3 days and 3 nights of measurements from patient 11 were excluded because correct time stamps were missing, due to the patient unintentionally resetting the time indication by taking out the batteries. The last day and night of patient 11’s measurements could not be analyzed because no clear distinction could be made between daytime and nocturnal data. Still, 3 days and 3 nights of patient 11’s measurements were used for further analyses.

The SpO$_2$ dataset contained 1.83% error values and an additional 1.49% were excluded during the first two steps of the data-cleaning algorithm (see Figure 3), resulting in 96.68% of valid data. Nocturnal data had 99.31% of valid data (0.45% error data and 0.24% cleaned) compared to 93.27% of valid data during the day (3.62% error data and 3.11% cleaned). This was similar when only considering daytime data during rest and LIPA (EE$\leq$3 MET; 93.90% of valid data: 3.25% error data and 2.85% cleaned). However, during MVPA (EE$>$3 MET), the amount of valid data decreased to 67.86% (18.34% error data and 13.80% cleaned).
Table 2. Summary of the amount of peripheral blood oxygen saturation (SpO₂) measurements that were included for further analyses.a.

<table>
<thead>
<tr>
<th>Measurements included for further analyses</th>
<th>Mean (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Days per patient</td>
<td>5.3 (1.8)</td>
</tr>
<tr>
<td>Nights per patient</td>
<td>5.9 (1.2)</td>
</tr>
<tr>
<td><strong>Hours per day</strong></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>7.8 (3.9)</td>
</tr>
<tr>
<td>During rest ($\text{EE}^{b} \leq 1.5 \text{ MET}^{c}$)</td>
<td>6.1 (3.1)</td>
</tr>
<tr>
<td>During LIPA$^{d}$ ($1.5 \text{ MET} &lt; \text{EE} \leq 3 \text{ MET}$)</td>
<td>1.5 (2.7)</td>
</tr>
<tr>
<td>During MVPA$^{e}$ ($\text{EE} &gt; 3 \text{ MET}$)</td>
<td>0.2 (0.2)</td>
</tr>
<tr>
<td><strong>Hours per night</strong></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>8.0 (0.9)</td>
</tr>
<tr>
<td>During the day</td>
<td>296,533 (95,580)</td>
</tr>
<tr>
<td>During the night</td>
<td>149,665 (88,909)</td>
</tr>
<tr>
<td>All 20 patients performed nocturnal measures and 17 patients performed daytime measurements. Therefore, averages were calculated with 17 patients for daytime measurements and 20 patients for nocturnal and total measurements.</td>
<td></td>
</tr>
<tr>
<td><strong>Data samples per patient</strong></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>169,318 (35,488)</td>
</tr>
<tr>
<td>During the day</td>
<td>149,665 (88,909)</td>
</tr>
</tbody>
</table>

aAll 20 patients performed nocturnal measurements and 17 patients performed daytime measurements. Therefore, averages were calculated with 17 patients for daytime measurements and 20 patients for nocturnal and total measurements.

bEE: energy expenditure.

cMET: metabolic equivalent of task.

dLIPA: low-intensity physical activity.

eMVPA: moderate-to-vigorous physical activity.

Figure 3. The amount of invalid data of continuous peripheral blood oxygen saturation (SpO₂) measurements for different activities. LIPA: low-intensity physical activity; MVPA: moderate-to-vigorous physical activity.

SpO₂ Features and Fluctuations

The weekly averages of mean SpO₂ were 89.9% (SD 3.4), 91.9% (SD 3.1), and 92.1% (SD 2.9) for nocturnal (n=20), daytime (n=17), and daytime-in-rest (n=17) measurements, respectively. Figure 4 shows that mean nocturnal SpO₂ was lower than mean daytime SpO₂ in rest ($P < 0.001$). Weekly averages of mean SpO₂ were between 84.7% and 96.0% for daytime SpO₂ in rest and between 80.3% and 94.3% for nocturnal SpO₂ (see Figure 4).

The weekly averages of CT90 were 35% (SD 34), 20% (SD 29), and 18% (SD 29) for nocturnal, daytime, and daytime-in-rest measurements, respectively. No significant correlations were found between patient characteristics and weekly averages of the SpO₂ features.

Intraday and intranight SpO₂ fluctuations were quantified as the weekly average of SpO₂ SD for nocturnal (1.6%, SD 0.6), daytime (1.8%, SD 0.7), and daytime-in-rest (1.6%, SD 0.6) measurements. Nocturnal, daytime, and daytime-in-rest SpO₂...
SD values were inversely correlated with nocturnal, daytime, and daytime-in-rest mean SpO\textsubscript{2} values (all: \(P<.001\), \(R^2>0.4\)), respectively. On average, SpO\textsubscript{2} in rest values ranged over 10.8\% (SD 4.4) within one day. There was a significant difference between mean SpO\textsubscript{2} in the afternoon and in the evening \((P<.001)\) (see Multimedia Appendix 3), with an average difference of 0.8\% (SD 0.7) for those days where patients performed at least one hour of measurements during both the afternoon and evening. For this comparison, a total of 46 days from 12 different patients were used (median of 4 days per patient).

Night-to-night and day-to-day SpO\textsubscript{2} fluctuations were quantified as the weekly ranges of mean SpO\textsubscript{2} (2.0\%, SD 1.1; 1.9\%, SD 1.5; 1.9\%, SD 1.5) and CT90 (28\%, SD 25; 19\%, SD 20; 18\%, SD 21) for nocturnal, daytime, and daytime-in-rest measurements, respectively. A more detailed analysis of the night-to-night changes in CT90 is shown in Figure 5. When considering the definition of a nocturnal desaturator, three types of patients can be distinguished: consistent desaturators (5/20, 25\%), consistent nondesaturators (5/20, 25\%), and occasional desaturators that changed category over the 7 nights (10/20, 50\%). A total of 6 occasional desaturators (6/10, 60\%) were desaturators or nondesaturators in the first two nights, but changed category when SpO\textsubscript{2} measurements of another night were considered. The weekly average of mean SpO\textsubscript{2} for occasional desaturators was between 89.5\% and 91.9\%, compared to 80.3\%-88.8\% and 91.8\%-94.3\% for consistent desaturators and consistent nondesaturators, respectively.
Figure 4. Mean nocturnal peripheral blood oxygen saturation (SpO₂) compared to mean daytime SpO₂ in rest. The dots indicate weekly averages of mean SpO₂ for every patient, lines indicate the standard deviation of the mean SpO₂ values over different days and nights, and the orange line is the line of equality.
Figure 5. Cumulative time spent with peripheral blood oxygen saturation (SpO₂) below 90% (CT90) values for every night of every patient showed that 50% (10/20) of the included patients changed category between desaturator and nondesaturator. The number next to each dot indicates the corresponding night of the measuring week and the orange line indicates the threshold of CT90 (30%) that divides nights with and without desaturation. A missing night number indicates that no data was available for that night.

Discussion

Continuous SpO₂ measurements with wearable pulse oximeters identified significant SpO₂ fluctuations between and within multiple days and nights of patients with COPD. COPD patients are known to be a heterogeneous group [32,33], which leads to large differences in mean SpO₂ values between patients. Day-to-day SpO₂ fluctuations have not yet received a lot of attention in scientific literature and night-to-night fluctuations have not yet been studied over more than two nights in patients with COPD. Some studies examined intraday fluctuations, however, these studies only examined short-term SpO₂ fluctuations (ie, measurements of one hour or less) in healthy individuals [29] or infants [26-28]. Our results showed that significant SpO₂ fluctuations occurred between and within multiple days and nights for the included patients. This is often not taken into account in telemonitoring applications [12]. When using the current definition of nocturnal desaturation, 50% (10/20) of the included patients changed category between desaturator and nondesaturator over the course of 1 week.

Continuous SpO₂ measurements with high amounts of valid data could be obtained during a 7-day monitoring period, except during MVPA (EE>3 MET). Nocturnal measurements were well-perceived by the patients, leading to the high compliance. These measurements only had a small amount of data that was considered invalid (0.69%). This indicates that it is feasible to perform multi-night continuous SpO₂ measurements. Most patients also performed daytime measurements during rest and LIPA (ie, 17 patients measured, on average, 7.6 hours per day). These had a limited amount of invalid data (6.10%). However, very few measurements were performed during MVPA (ie, 0.2 hours per day) and almost one-third of these measurements produced invalid data due to motion artifacts. Therefore, it might not be recommended to analyze continuous SpO₂ measurements during periods of MVPA.

Two types of preprocessing steps for continuous SpO₂ data have been reported: (1) removing deviating samples or (2) down-sampling the data. Deviating samples have been removed visually [29,34] by using an unspecified data-cleaning algorithm [35] or by removing samples with a sample-to-sample deviation of more than 4% [36] or 8% [37]. The latter method, however, does not remove artifacts that last over longer periods of time (eg, see Figure 2B: around 17:00-18:00). Therefore, artifacts in this study were removed by removing blocks of deviating values instead of separate samples. After sample removal, Morillo et al excluded remaining artifacts by down-sampling the data from 8 Hz to 1 Hz [37]. However, autocorrelation analyses in our study showed that a sampling time of 1 sample per second still led to oversampling of the data (see Multimedia Appendix 2). SpO₂ could be down-sampled to 1 sample per 20 seconds for improved artifact removal, without losing information about the SpO₂ dynamics.

Mean nocturnal SpO₂ was lower than mean daytime SpO₂ in rest. By taking into account the SpO₂ fluctuations over different days and nights, the results of this study generalize the findings of Soguel Schenkel et al, who only performed SpO₂ measurements during a single day and night that were up to five days apart [38]. Other studies used awake measurements of
SpO₂ to predict nocturnal mean SpO₂ or CT90. They used the first 15-30 minutes of the nocturnal measurements to calculate the mean awake SpO₂ values [15,39-41]. For these studies, it should be noted that slightly lower SpO₂ values were observed in the evening compared to the afternoon, on average a 0.8% difference, and awake SpO₂ values depend on the time of day when the measurements are performed. It is unlikely that this difference is solely due to the specified oximeter accuracy of ±2%, as no consistent bias has been reported for the WristO₂ when measuring over a longer time period.

The observed patient-to-patient, day-to-day, and intraday differences can have implications for telemonitoring applications that are based on daily SpO₂ spot checks [12]. In several telemonitoring setups, alerts were raised when daily spot checks of SpO₂ dropped below a generic threshold value (eg, 90% for all patients) [12]. However, the high patient-to-patient differences point out the shortcomings of these generic thresholds. Other telemonitoring setups used personalized thresholds, but these were still fixed on one specific threshold value for every patient [12]. For these setups, alerts can be merely a consequence of natural day-to-day and intraday fluctuations, instead of being triggered by the onset of an exacerbation. A recent paper better dealt with the day-to-day fluctuations by suggesting a day-to-day decrease of more than 4% to alert for exacerbations, however, a more thorough examination of this method is needed [42]. Intraday SpO₂ values during rest ranged over 10.8% (SD 4.4) within one day. Moreover, statistical interpretation of weekly average of SpO₂ SD (1.6%, SD 0.6), which is a measure of intraday fluctuations, indicates that during 5% of the day in rest, SpO₂ fluctuates more than 3.2% (ie, 2 times SpO₂ SD) beyond the mean daytime-in-rest value. In comparison, previous studies only reported a 1%-2% decrease in SpO₂ spot checks around exacerbation onset compared to stable periods [43-45]. These natural intraday fluctuations can thus easily result in false alerts for exacerbations. Patients with lower SpO₂ values might experience a higher number of false alerts as, similar to healthy individuals [29], intraday fluctuations (ie, SpO₂ SD) increased with decreasing mean SpO₂. Altogether, personalized alerts based on intelligent algorithms will be necessary to cope with all of these natural fluctuations in daytime SpO₂ in rest. Preferably, these alerts should be based on continuous measurements over longer time periods, in contrast to the currently used daily spot checks, to account for the identified SpO₂ fluctuations and to exclude the potential effect of the ±2% oximeter accuracy. In addition, the slightly lower SpO₂ values in the evening compared to the afternoon show that the daily measurements should always be performed at the same moment of the day. A clearly defined measurement protocol, which is often not specified [12], can thus further attempt to limit the effect of natural SpO₂ fluctuations in these telemonitoring applications.

The night-to-night differences of nocturnal mean SpO₂ (ie, average weekly range of 2.0% over the different nights) led to highly varying CT90 values over the different nights of the measurement week (ie, average range of 28%). These high variations in CT90 resulted in 50% (10/20) of the included patients changing category between desaturator and nondesaturator, due to the fact that these occasional desaturators all had a mean SpO₂ value around the threshold of 90%. A similar finding has been reported by Lewis et al, who concluded that 35% of the included patients changed category over two consecutive nights of measurements [20]. Later studies then tackled this problem by performing measurements over two nights, categorizing a patient as a desaturator if desaturation occurred in at least one of both nights [5,17-19]. However, the results of this study indicate that even two nights are insufficient to make a consistent separation between desaturators and nondesaturators, as 6 out of 10 occasional desaturators (60%) were desaturators or nondesaturators in the first two nights and only changed category afterward. Our results suggest that it might be impossible to consistently categorize COPD patients with a mean nocturnal SpO₂ value around 90% as desaturator or nondesaturator. Based on measurements over multiple nights, these patients could thus be referred to as occasional desaturators. Further research is needed to assess the clinical relevance of identifying these three different nocturnal desaturation profiles.

Some limitations should be taken into consideration when interpreting the results of this study. The main limitation, as is often the case in similar studies, is the small sample size. Consequently, no comparison could be made between mild, moderate, and severe hypoxemic patients. Nevertheless, the increasing SpO₂ SD with decreasing mean SpO₂ suggests an increase in SpO₂ fluctuations for more hypoxemic patients. No control group was included because this study aimed to perform continuous SpO₂ measurements in COPD patients for identification of SpO₂ fluctuations that could affect SpO₂ applications, rather than comparing SpO₂ between COPD patients and healthy controls. In addition, daily home routines can greatly differ between patients and healthy controls [46], impeding a proper comparison. As shown in Multimedia Appendix 1, daytime measurements were only seldom performed during full days due to the impracticalities of wearing the finger clip during activities that require manual actions (eg, during morning routines). This, in combination with the high amount of invalid data during MVPA, suggests that continuous SpO₂ measurements with a finger clip will have more profound limitations in a population that is more physically active than the target population of this study. Reliable wearable oximeters that do not require a finger clip could thus increase compliance. Moreover, the limited battery life and the inability of real-time data transmission of the used oximeter can further complicate the integration of continuous SpO₂ measurements into practice. Technological advances are thus needed to allow for long-term, continuous monitoring of SpO₂. This study, however, mainly intended to show the potential of prolonged continuous measurements to identify SpO₂ fluctuations. Therefore, a certified wearable pulse oximeter with finger clip was preferred over more user-friendly, watch-type oximeters that have not yet been proven to be accurate. The resulting, more fragmented, daytime measurements were sufficient to identify large natural
fluctuations occurring within one day (ie, SpO\textsubscript{2} in rest ranged over 10.8% [SD 4.4] within one day), confirming the a priori posed hypothesis. The limited battery life was addressed by providing the patients with multiple sensors to cover the 7-day monitoring period.

This study was the first to use wearable finger pulse oximeters for prolonged continuous SpO\textsubscript{2} measurements in COPD patients, as opposed to only performing spot checks or continuous measurements during one or two nights. These measurements showed that spot checks or one- or two-night measurements should be interpreted with caution, as the conclusions based on these measurements might change depending on the moment of the measurement. Measurements were performed at home during daily routine COPD patients, which provides a more natural SpO\textsubscript{2} profile compared to supervised measurements. By adding actigraphy measurements, the necessary contextual information could be gathered for more accurate analyses of the continuous SpO\textsubscript{2} measurements.

In conclusion, continuous SpO\textsubscript{2} measurements with wearable pulse oximeters identified significant SpO\textsubscript{2} fluctuations between and within multiple days and nights of patients with COPD. Continuous SpO\textsubscript{2} measurements during the daily home routine of patients with COPD generally had high amounts of valid data, except for motion artifacts during MVPA. The continuous measurements showed that mean nocturnal SpO\textsubscript{2} was lower than mean daytime SpO\textsubscript{2} in rest, and significant SpO\textsubscript{2} fluctuations occurred between and within multiple days and nights. The large fluctuations of daytime SpO\textsubscript{2} in rest indicate that clear measurement protocols and personalized alerts, based on intelligent algorithms, will be needed to increase the performance of telemonitoring applications that make use of daily SpO\textsubscript{2} spot checks. Lastly, it was shown that CT90 values can vary greatly from night to night in patients with a nocturnal mean SpO\textsubscript{2} around 90%, indicating that these patients cannot be consistently categorized as desaturators or non-desaturators. We recommend using wearable sensors for performing continuous SpO\textsubscript{2} measurements over longer time periods to determine the clinical relevance of the identified SpO\textsubscript{2} fluctuations.

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Conflicts of Interest
MAS discloses receipt of renumeration for consultancy and/or lectures from Boehringer Ingelheim, GlaxoSmithKline, and AstraZeneca outside the scope of this work. The authors have no other relevant affiliations or financial involvement with any organization or entity with a financial interest in or financial conflict with the subject matter or materials discussed in the manuscript, apart from those disclosed.

Multimedia Appendix 1
Visual overview of the continuous peripheral blood oxygen saturation (SpO\textsubscript{2}) measurements that were included for analyses.

[PDF File (Adobe PDF File), 32KB-Multimedia Appendix 1]

Multimedia Appendix 2
Autocorrelation analyses.

[PDF File (Adobe PDF File), 95KB-Multimedia Appendix 2]

Multimedia Appendix 3
Differences between mean peripheral blood oxygen saturation (SpO\textsubscript{2}) in the afternoon and in the evening. Grey dots indicate the differences within the same day for the same patient and orange dots indicate the average of these differences for every patient. The number of grey dots per patient indicate the amount of days with measurements in both the afternoon and evening.

[PDF File (Adobe PDF File), 22KB-Multimedia Appendix 3]

References


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**Abbreviations**

- **6MWD**: 6-minute walking distance
- **CAT**: chronic obstructive pulmonary disease assessment test
- **CIRO**: Centre of Expertise for Chronic Organ Failure
- **COPD**: chronic obstructive pulmonary disease
- **CT90**: cumulative time spent with SpO₂ below 90%
- **EE**: energy expenditure
- **FEV1**: forced expiratory volume in 1 second
- **FVC**: forced vital capacity
- **GOLD II-IV**: Global Initiative for Chronic Obstructive Lung Disease stages II-IV
- **ISO**: International Organization for Standardization
- **LIPA**: low-intensity physical activities
- **MEC-U**: Medical Research Ethics Committees United
- **MET**: metabolic equivalent of task
- **MVPA**: moderate-to-vigorous physical activities
- **PaO₂**: oxygen partial pressure in arterial blood
- **Seff**: sleep efficiency
- **SpO₂**: peripheral blood oxygen saturation
- **TLCO**: transfer factor for carbon monoxide
- **TNST**: total night sleeping time
- **VITO**: Flemish Institute for Technological Research
- **WASO**: wake time after sleep onset

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