



PULMONOLOGY

www.journalpulmonology.org


LETTER TO THE EDITOR

Use of automatic 6-minute walking test recording system in patients with chronic respiratory diseases

KEYWORDS

6MWT;
Distance sensors;
Biological signals;
COPD;
Interstitial lung disease;
Pulmonary hypertension

To the editor:

Physical capacity tests are used for the diagnosis, prognosis and monitoring of chronic respiratory diseases (CRD), such as Chronic Obstructive Pulmonary Disease (COPD) or Pulmonary Hypertension (PH).¹ Among these tests, the most widely used is the 6-minute walking test (6MWT).² Additionally, the guidelines highlight the need to continuously record biological signals, particularly oxygen saturation (SpO₂), which has been shown to be a prognostic marker in CRD.²

Technological advances have made available new solutions that improve monitoring and testing, including the 6MWT.³ By applying off-the-shelf technology, including the internet of things (IoT), edge computing and wireless communications, the required biological signals can be automated and recorded in a more user-friendly and reliable way, thus meeting recommendations precisely.⁴ Our objective was to determine the reliability and agreement in capturing the SpO₂ and heart rate (HR) data of an automated 6MWT recording system in patients with COPD or PH, in comparison with that manually recorded by a technician.

We conducted a cross-sectional study of patients with a COPD or PH diagnosis followed at the Hospital Clínic in Barcelona, Spain, between October 2022 and March 2023. Ethics committee approval was obtained, and all patients signed informed consent forms. The inclusion criteria were patients older than 18 years and diagnosis of COPD or PH. Exclusion criteria were the presence of locomotor or cognitive impairment and any pre-existing condition limiting the ability to perform at test. Anthropometric characteristics, pulmonary diagnostics, and lung function parameters were

collected. The main outcomes were SpO₂ and HR during the 6MWT.

The 6MWT was conducted indoors in a flat, straight, 30-metre walking corridor.² The modified Borg scale (0–10) was used to measure dyspnoea and fatigue.⁵ A finger oximeter (Nonin WristOx 3130, Plymouth, USA) was used to record SpO₂ and HR. We collected patients' data for an analysis corresponding to 8 data points (baseline, final and every minute during the test).

The 6MWT automatic data collection system consists of a non-commercial prototype designed in a doughnut shape for placement in any type of cone. To record subjects' counter-clockwise turns around the cone, it uses sensors that detect movement and send data to a tablet with an ad-hoc application. Simultaneously, the application records HR and SpO₂ biological signals that are acquired at 1 Hz frequency through the finger oximeter. First, the technician must set up the scenario, place sensors over the cones and attach the pulse oximeter to the patient. Meanwhile, the two sensors and the pulse oximeter are paired to the application. The test starts counting the elapsed time when the patient turns around the cone. Then, each time the patient turns around a checkpoint, a card will appear in the application, displaying the test's elapsed time, the time difference from the previous card, the checkpoint number, the SpO₂, and the HR. At the end of the test, the system automatically computes the extra meters walked from the last checkpoint and asks the patient again about dyspnoea and fatigue. When the test is finished, all raw data obtained from the pulse oximeter in all test stages (including the initial rest and the final recuperation) are uploaded, highlighting the checkpoint data. It is accessible to anyone with the required authorisation. A trained technician manually recorded the biological signals in an ad-hoc form. As in the automatic system, the variables were recorded each time the patient passed through one of the cones.

Using the method of Walter et al.⁶ based on an estimate using the intraclass correlation coefficient (ICC), at least 40 individuals are necessary, considering an acceptable reliability of $p_0 = 0.60$ and an expected reliability of $p_1 = 0.80$, with a power of 90 % and a level of significance of 5 %, due to the nature and characteristics of the study, a loss of 5 % of the sample is assumed. All data are expressed as means, standard deviations (SD) or medians at the 25th and 75th percentile depending on the distribution. The distribution was analysed using the Shapiro-Wilk test. Differences between

<https://doi.org/10.1016/j.pulmoe.2023.08.011>

2531-0437/© 2023 Sociedade Portuguesa de Pneumologia. Published by Elsevier España, S.L.U. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

Please cite this article in press as: R. Torres-Castro, H. Pascual, A. Alonso et al., Use of automatic 6-minute walking test recording system in patients with chronic respiratory diseases, Pulmonology (2023), <https://doi.org/10.1016/j.pulmoe.2023.08.011>

Table 1 Descriptive data of the included patients.

Variable	N = 45	
Sex, M/F	9/36	
Age, years	58 ± 15	
Weight, Kg	68 ± 15	
Height, cm	164 ± 9	
BMI, Kg/m ²	25.0 ± 4.9	
<i>Diagnosis</i>		
PH	27 (60.0 %)	
COPD	18 (40.0 %)	
<i>Pulmonary function</i>	COPD	PH
FEV ₁ , % ref	50 ± 14	84 ± 17
FVC, % ref	82 ± 17	93 ± 14
FEV ₁ /FVC, %	45 ± 10	71 ± 10
DL _{CO} , % ref	57 ± 18	57 ± 16
K _{CO} , % ref	60 ± 15	65 ± 19
<i>Six-minute walking test</i>		
Distance walked, metres	493 ± 121	
Dyspnoea baseline, Borg	1 (0-2)	
Dyspnoea at the end, Borg	3 (2-4)	
Fatigue baseline, Borg	1 (0-1)	
Fatigue at the end, Borg	2 (1-3)	
<i>Heart rate*</i>		
Manual system baseline, beat/min	85 ± 17	
Automatic system baseline, beat/min	85 ± 18	
Manual system at the end, beat/min	116 ± 19	
Automatic system at the end, beat/min	112 ± 21	
<i>Oxygen saturation*</i>		
Manual system baseline, %	96.4 ± 1.6	
Automatic system baseline, %	96.3 ± 1.6	
Manual system at the end, %	89.4 ± 7.2	
Automatic system at the end, %	89.3 ± 6.8	

Values are expressed as the mean ± SD if data are normally distributed or as the median (P25–P75) if the data distribution is skewed. *The difference between both systems > 0.05. Definition of abbreviations: BMI: Body mass index; PH: Pulmonary hypertension; COPD: Chronic obstructive pulmonary disease; FEV₁: Forced expiratory volume in the first second; FVC: Forced vital capacity; DL_{CO}: Diffusing capacity for carbon monoxide; K_{CO}: Carbon monoxide transfer coefficient.

data were evaluated using Student's t-test for normally distributed variables or a Mann–Whitney U test for non-parametric variables. The ICC and Bland–Altman plots were used to evaluate the agreement between both systems. All statistical analyses were performed with the SPSS Version 25.0 package (SPSS, Chicago, IL, USA).

We recruited 45 patients (36 [80 %] women; 58 ± 15 years). The baseline characteristics of the patients are presented in Table 1. Regarding the 6MWT, the mean distance walked (in the manual and automatic registry) was 493 ± 121 m. The ICC for the HR was 0.894 (0.863–0.918) and for SpO₂, it was 0.955 (0.941–0.965). The Bland–Altman plot showed that 94.2 % of the HR data and 94.2 % of the SpO₂ data were within the limits of agreement (Fig. 1).

The automatic system can acquire biological signals at a frequency of 1 Hz, thus providing much more information during the performance of the 6 MWT. In contrast, when it is manually recorded, biological signals usually cannot be collected due to the limitations of technicians who first need to pay attention to patients' safety.

Finally, to date, automatic recording systems have been developed that mainly focus on remote evaluation or the recording of biomechanical gait parameters.³ The proposed system, unlike those previously published, focuses on the recording of biological signals in respiratory patients, especially SpO₂, which predict mortality and exacerbation.^{7,8}

In conclusion, using an automated recording system based on distance sensors and the transmission of biological signals is reliable and shows excellent agreement with the manual registration system for the clinical registration of SpO₂ and HR during the 6MWT in patients with CRD.

Conflict of interest

The authors declare to have no conflict of interest.

CRedit authorship contribution statement

R. Torres-Castro: Formal analysis, Methodology, Writing – original draft, Writing – review & editing. **H. Pascual:** Formal analysis, Methodology, Writing – review & editing. **A. Alonso:** Conceptualization, Formal analysis, Methodology, Writing – review & editing. **E. Gimeno-Santos:** Methodology, Writing – review & editing. **J.A. Barberà:** Formal

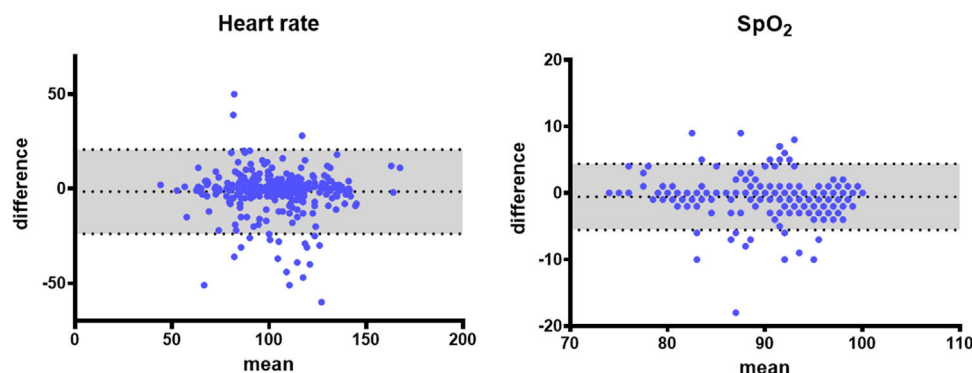


Fig. 1 Bland–Altman plots for biological signals.

analysis, Methodology, Writing – review & editing. **J. Bigorra**: Writing – review & editing. **J. Batlle**: Conceptualization, Writing – review & editing. **X. Masip-Bruin**: Conceptualization, Formal analysis, Methodology, Writing – review & editing. **I. Blanco**: Conceptualization, Formal analysis, Writing – original draft, Writing – review & editing.

Funding

The study was supported by grants from Societat Catalana de Pneumologia (SOCAP) (ESTEVE TEIJIN/2021), [PI17/1515](#) and [PI21/0555](#) from the Instituto de Salud Carlos III (ISCIII), co-funded by the European Union (ERDF/ESF, “A way to make Europe” “Investing in your future”) and Premi d’Innovació de l’Hospital Clínic de Barcelona (2021). For UPC authors this work has been supported by the Spanish Ministry of Science and Innovation under contract [PID2021-124463OB-I00](#), the Catalan Government under contract 2021 [SGR 00326](#) and the Catalan Department of Research and Universities.

Acknowledgments

Rodrigo Torres-Castro is a Ph.D. candidate in Methodology of Biomedical Research and Public Health Program, at Universitat Autònoma de Barcelona, Barcelona, Spain.

References

1. Clini EM, Crisafulli E. Exercise capacity as a pulmonary rehabilitation outcome. *Respiration*. 2009;77:121–8.
 2. Holland AE, Spruit MA, Troosters T, Puhan MA, Pepin V, Saey D, et al. An official European respiratory society/American thoracic society technical standard: Field walking tests in chronic respiratory disease. *Eur Respir J*. 2014;44:1428–46.
 3. Pires IM, Denysyuk HV, Villasana MV, Sá J, Marques DL, Morgado JF, et al. Development technologies for the monitoring of six-minute walk test: a systematic review. *Sensors*. 2022;22:581.
 4. Angelucci A, Aliverti A. Telemonitoring systems for respiratory patients: technological aspects. *Pulmonology*. 2020;26:221–32.
 5. Johnson MJ, Close L, Gillon SC, Molassiotis A, Lee PH, Farquhar MC, et al. Use of the modified Borg scale and numerical rating scale to measure chronic breathlessness: a pooled data analysis. *Eur Respir J*. 2016;47:1861–4.
 6. Walter SD, Eliasziw M, Donner A. Sample size and optimal designs for reliability studies. *Stat Med*. 1998;17:101–10.
 7. Liu S-F, Chin C-H, Tseng C-W, Chen Y-C, Kuo H-C. Exertional desaturation has higher mortality than non-desaturation in COPD. *Medicina (B. Aires)*. 2021;57:1110.
 8. Vainshelboim B, Kramer MR, Izhakian S, Lima RM, Oliveira J. Physical activity and exertional desaturation are associated with mortality in idiopathic pulmonary fibrosis. *J Clin Med*. 2016;5:73.
- R. Torres-Castro^{a,b,c}, H. Pascual^d, A. Alonso^b, E. Gimeno-Santos^{a,e,f}, M. Palomo^a, J.A. Barberà^{a,b,f}, J. Bigorra^{b,e}, J. Batlle^g, X. Masip-Bruin^d, I. Blanco^{a,b,f,*}
- ^a Department of Pulmonary Medicine, Hospital Clínic, University of Barcelona, Barcelona, Spain
^b Fundació de Recerca Clínic Barcelona-Institut d’Investigacions Biomèdiques August Pi i Sunyer (IDIBAPS), Barcelona, Spain
^c Department of Physical Therapy, Faculty of Medicine, University of Chile, Santiago, Chile
^d CRAAX Lab, Universitat Politècnica de Catalunya, Vilanova i la Geltru, Spain
^e Instituto de Salud Global (ISGlobal), Barcelona, Spain
^f Biomedical Research Networking Center on Respiratory Diseases (CIBERES); Madrid, Spain
^g Innovation Consultant-Techno & Innovative Solutions, SLU; Vilanova i la Geltru, Spain
- * Corresponding author at: Department of Pulmonary Medicine, Hospital Clínic-IDIBAPS, University of Barcelona, Barcelona, Spain.
 E-mail address: iblanco2@clinic.cat (I. Blanco).
 Received 20 April 2023; Accepted 31 August 2023
 Available online xxx