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Variability in objective and subjective measures affects baseline values in studies of patients with COPD

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Abstract

Rationale

Understanding the reliability and repeatability of clinical measurements used in the diagnosis, treatment and monitoring of disease progression is of critical importance across all disciplines of clinical practice and in clinical trials to assess therapeutic efficacy and safety.



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Competing interests: This repeatability study is a sub-study of SPIROMICS. SPIROMICS is a multicenter observational study of COPD designed to guide future development of therapies for COPD The design of SPIROMICS and this Substudy was approved and funded by the NIH through contracts to clinical sites. The NIH did not have a role in the analysis, decision to publish or the preparation of this manuscript. Support for the study was also provided by the Foundation for the NIH (FNIH). The FNIH procures funding and manages alliances with public and private institutions in support of the mission of the National Institutes of Health (NIH). No funds were received directly from the Companies listed in the funding source. These Companies had no role in study design, data collection, analysis, the decision to publish or preparation of this manuscript. Dr. Rennard was Professor of Medicine at the University of Nebraska at the time this study was approved and funded. Dr. Cooper is now a part-time employee of GlaxoSmithKline. None of the other authors are employees of these companies, nor received compensation related to this study.

Objectives

Our goal is to understand normal variability for assessing true changes in health status and to more accurately utilize this data to differentiate disease characteristics and outcomes.

Methods

Our study is the first study designed entirely to establish the repeatability of a large number of instruments utilized for the clinical assessment of COPD in the same subjects over the same period. We utilized SPIROMICS participants (n = 98) that returned to their clinical center within 6 weeks of their baseline visit to repeat complete baseline assessments. Demographics, spirometry, questionnaires, complete blood cell counts (CBC), medical history, and emphysema status by computerized tomography (CT) imaging were obtained.

Results

Pulmonary function tests (PFTs) were highly repeatable (ICC's >0.9) but the 6 minute walk (6MW) was less so (ICC = 0.79). Among questionnaires, the Saint George's Respiratory Questionnaire (SGRQ) was most repeatable. Self-reported clinical features, such as exacerbation history, and features of chronic bronchitis, often produced kappa values <0.6. Reported age at starting smoking and average number of cigarettes smoked were modestly repeatable (kappa = 0.76 and 0.79). Complete blood counts (CBC) variables produced intraclass correlation coefficients (ICC) values between 0.6 and 0.8.

Conclusions

PFTs were highly repeatable, while subjective measures and subject recall were more variable. Analyses using features with poor repeatability could lead to misclassification and outcome errors. Hence, care should be taken when interpreting change in clinical features based on measures with low repeatability. Efforts to improve repeatability of key clinical features such as exacerbation history and chronic bronchitis are warranted.

Introduction

The Subpopulations and Intermediate Outcome Measures in COPD Study (SPIROMICS) is an observational study of 2,981 participants including: healthy never-smokers, ever-smokers (> 20 pack/years) with preserved PFTs, and individuals with COPD classified as mild-moderate or severe by PFTs designed to aid in the future development of therapies for COPD [1]. The Repeatability Substudy embedded in SPIROMICS consisted of 98 participants who volunteered to return within 2–6 weeks to repeat their baseline visit. We designed this Substudy to quantify baseline, within-person variation, including measurement errors. The 2–6 week window of the SPIROMICS Repeatability Substudy is considered short enough to avoid changes due to disease progression, yet long enough to minimize any learning effect from the initial visit. Assessing the severity of COPD utilizes a number of objective (PFTs, 6-min walk distance), subjective [modified medical research council (mMRC) dyspnea, COPD assessment test (CAT)] and patient recall (number of exacerbations) measures utilized in determining disease progression, risk of exacerbations and treatment effects. Consistency of clinical assessments during periods of disease stability is critical to interpretation, but often goes unreported. This report adds to recent reports on functional tests in COPD [2] by describing the repeatability of the selected assessments and their potential impact on assessments of COPD used in classifying severity, stability and progression. Some of the results of this study have been previously reported in the form of an abstract [3].

Methods

SPIROMICS is a prospective cohort study that enrolled 2981 participants at 11 clinical sites. Extensive coordinator training, monitoring and follow-up was provided to assure consistency across all sites. All subjects were age 40 to 80 years and willing to undergo the extensive study procedures. Exclusion criteria included BMI >40 kg/m², cognitive dysfunction and other lung disease or clinically significant cardiovascular disease that would limit the interpretability of the measures. A history of asthma was exclusionary only for never-smokers. The full SPIRO-MICS study design and inclusion and exclusion criteria have been reported [1], and the Repeatability dataset is available upon request (www.spiromics.org).

The SPIROMICS Repeatability Substudy comprised 98 subjects from the largest clinical sites, who repeated their entire baseline evaluation 2-6 weeks after their initial visit. Subjects experiencing an exacerbation between the baseline and repeat study visits were excluded from this analysis. We separated the evaluations into A) objective measures [PFTs, six minute walk distance and CBC], B) measures dependent on patient recall [medical history including previous lung disease diagnosis, smoking history and exacerbations] and C) subjective evaluations [CAT[4], SGRQ [5], Functional Assessment of Chronic Illness Therapy-Fatigue [6] (FACIT-F) Score, Pittsburg Sleep Quality Index [7] (PSQ), and Medical Outcomes SF-12 [8]]. PFTs were conducted pre- and post- bronchodilation (BD) (4 puffs of Albuterol sulfate HFA plus 4 puffs of Ipratropium bromide HFA). Withholding bronchodilators was not required; time from last administration was recorded. We evaluated the repeatability of a diagnosis of chronic bronchitis (CB) using: 1) patient recall of being diagnosed with CB by a health care professional; or 2) the classic definition of chronic cough and sputum production for at least 3 months/year for the last two consecutive years [9]; and 3) from the SGRQ questions regarding cough and phlegm (cough and phlegm production several days a week or almost every day)[10]. Emphysema was assessed by 1) recall of a diagnosis by a health care professional and 2) diagnosed from volumetric multidetector-row computed tomography (MDCT) of the lungs performed at full inspiration, using an emphysema index (EI) of percent voxels in the lung field < -950 HU. GOLD staging was calculated as reported [11]. Questionnaires were administered by a study coordinator in person, and answered solely by the participant. All blood samples for CBC differentials were prepared according to a standard SPIROMICS protocol and analyzed locally at each University Laboratory. All subjects provided written informed consent, and the study was approved by the IRB at each participating site (See S1 Study Information for additional details).

Statistical analysis

To provide reliability measurements, we calculated intraclass correlation coefficients (ICC) and kappa statistics for quantitative and qualitative traits (MRC breathlessness scale and current smoking status), respectively. When a value for a trait was missing for the baseline or repeat visits, the subject was dropped from the analysis for that trait. Bland-Altman plots were used to visualize potential changes in reliability across the spectrum of quantitative traits. In some cases, reliability measures were calculated for the entire Substudy cohort and for a subset diagnosed with COPD based upon baseline PFTs.

Results

Demographics

The Substudy subjects were predominately white males (except for never-smokers) and non-obese. The two COPD groups were significantly older than those without airways obstruction (p = 0.0006) (Table 1). There were no significant differences between groups in BMI, reported pack-years smoking (excluding never-smokers) or time between visits (28.9 ± 9.4 days, mean ± SD); among the ever-smoker groups, fewer participants with COPD reported smoking currently. Pulmonary function differences reflect SPIROMICS enrollment groups. Bronchodilator response, as a percent change, increased with increased disease severity. However, the absolute change in forced expiratory volume in one second (FEV₁) was not significantly different between the strata with absolute changes in FEV₁ of 172.0 ±233.4, 121.9 ± 150.0, 215.2 ± 148.0 and 216.4 ± 167.2 for strata 1–4 respectively (p \ge 0.314). The increasing FEV₁ reversibility for stratum 3 and 4 as percent change results from the decreasing FEV₁ as disease progresses [12].

We compared demographics (age, BMI, height, weight, pack-years) and post-bronchodilator PFTs by groups within the repeatability population at baseline (n = 98) to the total SPIRO-MICS cohort (n = 2852; Repeatability Substudy subjects removed). There were no significant differences in pack-years or any of the demographic parameters, with the exception of age in

	Stratum 1	Stratum 2	Stratum 3	Stratum 4
	Never Smokers	Non-Diseased Smokers	Mild-Moderate COPD	Severe COPD
N	11	18	38	31
Age	55.4 ± 6.4	56.2 ± 8.4	64.7 ± 8.3	63.3±8.5
BMI	26.9 ± 4.5	29.5 ± 5.9	27.7 ± 5.1	26.6±5.9
Gender, % Male	36	61	76	61
Current Smokers %	0	72	54	30
Pack Years	0	41.2 ± 19.4	47.3 ± 19.6	47.5 ± 19.0
Race, %				
White	64	56	71	81
Black	18	28	21	13
Hispanic	18	5	3	6
Asian	0	6	0	0
American Indian	0	6	3	0
FVC L (% Predicted)	4.19 ± 1.30 (106.2)	3.87 ± 0.75 (97.5)	3.77 ± 1.0 (94.9)	2.92 ± 1.09 (72.7)
FEV ₁ L (% Predicted)	3.42 ± 1.04 (110.4)	2.99 ± 0.63 (97.7)	2.27 ± 0.71 (75.9)	1.14 ± 0.48 (38.1)
FEF _{25-75%} L/sec (% Predicted)	3.75 ± 1.16 (130.1)	2.84 ± 0.96 (101.0)	1.16 ± 0.56 (46.2)	0.46 ± 0.37 (18.0)
FEV ₁ /FVC (% Predicted)	103.6	100	79.4	52.9
Bronchodilator Response (% change) ⁺				
FEV ₁ (n)	5.5 ± 9.9 (10)	4.8 ± 5.8 (16)	12.6 ± 12.7 (25)	24.8 ± 24.2 (11)
FVC (n)	0.06 ± 6.0 (10)	0.81 ± 3.8 (16)	11.1 ± 10.3 (25)	18.0 ± 17.7 (11)
6-Min Walk Distance (M)	486.3 ± 46.6	426.5 ± 86.8	403.3 ± 111.3	329.3 ± 135.4

Table 1. Demographics, PFTs and six-minute walk.

Values are means ± SD; n = number of subjects used in the analysis. Stratum 1 = Non-Smokers (FVC > LLN and FEV1/FVC >0.7); Stratum 2 = Non-Diseased Smokers (FVC > LLN and FEV1/FVC >0.7); Stratum 3 = COPD FEV1 >50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FEV1 <50% and FEV1/FVC <0.7; Stratum 4 = COPD FE

⁺ Analysis was restricted to subjects who had not used any long-acting bronchodilator in the past 48 hours or a LABA within the past 24 hours, Tiotropium within the past 48 hours, a SABA within 6 hours or ipratropium within the last 8 hours of the baseline visit.

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Non-Diseased Smokers (p = 0.05) for whom mean ages were 56 vs. 60 years for the Repeatability Substudy and the overall cohort respectively. There was no difference between the Substudy and entire cohort in FEV₁ percent predicted.

Objective measures

PFTs were highly repeatable. A scatter plot and Bland-Altman plot for baseline and repeat visit post-bronchodilator FEV₁ are shown in Fig 1. Data for post-bronchodilator FVC, FEV₁/FVC and IC are presented in (S1 Fig). Paired differences (mean \pm SD) between visits were very small for post-bronchodilator FVC (0.04 \pm 0.34L), FEV₁ (0.02 \pm 0.19 L/sec), FEV₁/FVC (-0.001 \pm 0.04%), and IC (0.02 \pm 0.35 L). There was no apparent effect of severity on repeatability of these measures.

To reduce any potential impact of control subjects on the variability of PFTs, we calculated ICCs and the mean \pm SD values utilizing only COPD subjects with an FEV₁/FVC ratio <0.7 (**Table 2**). ICCs ranged from 0.81–0.97 for all measures, lowest for FVC and FEV₁ bronchodilator response. The mean value of the expiratory PFT measures on the repeat visit was slightly larger than the baseline values. The ICCs for PFTs by GOLD stage remained >0.92 for GOLD 1–3 but decreased to 0.82 for GOLD 4 subjects (n = 9) (data not shown). Repeatability (ICC) of absolute post- to pre- FEV₁ bronchodilator response for COPD subjects was 0.87 and FVC was 0.81.

The 6-minute walk distance (**Fig 2**) had lower repeatability (ICC = 0.79) compared to PFT measures, with a mean increased distance of 18.6 meters at the repeat visit, which is below the minimal clinically important difference (MCID) of 26 meters[13]. Subjects with the shorter walk distances were more variable between visits. Sixteen subjects (17.8%) had a decline greater than 26 meters and twenty-eight subjects (31.1%) had an increase of greater than 26 meters at the repeat visit. The mean MRC dyspnea score slightly improved from a mean of 1.5 to 1.3, consistent with the modest increase in 6-minute walk distance. Forty-one subjects

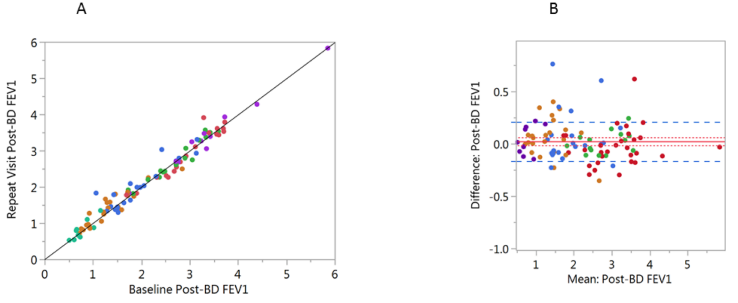


Fig 1. Scatterplot of baseline vs. repeat visit FEV₁. Subjects are color coded by GOLD stratification (using PFT values only). GOLD 0 = red, GOLD 1 = green, GOLD 2 = Blue, GOLD 3 = orange and GOLD 4 = Purple. The solid black line is drawn as a line of identity to visualize differences between baseline and the repeat visit. A) Correlation between the baseline and repeat visit FEV₁. (r = 0.983, p < 0.0001; n = 98). B) Bland-Altman plot of baseline mean post-bronchodilator FEV₁ by the difference between visits. The solid red line = the mean difference, the dotted red line is ± 1 SE and the dashed blue line is ± 1 SD.

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Clinical Measure	Baseline	Repeat Visit	ICC* COPD ⁺
Pre-Bronchodilator (Mean ± SD)			
FVC (I)	3.09 ± 1.10	3.17 ± 1.10	0.94
FEV ₁ (I)	1.57 ± 0.83	1.61 ± 0.81	0.97
FEF ₂₅₋₇₅ (l/sec)	0.73 ± 0.54	0.74 ± 0.56	0.91
PEFR (I/sec)	4.81 ± 2.24	4.95 ± 2.21	0.96
FEV ₁ /FVC	0.49 ± 0.14	0.50 ± 0.14	0.93
SVC (I)	3.19 ± 1.15	3.29 ± 1.11	0.95
IC (I)	2.44 ± 0.85	2.51 ± 0.85	0.94
Post-Bronchodilator (Mean ± SD)			
FVC (I)	3.39 ± 1.12	3.46 ± 1.08	0.95
FEV ₁ (I)	1.76 ± 0.84	1.81 ± 0.82	0.97
FEF ₂₅₋₇₅ (l/sec)	0.84 ± 0.59	0.85 ± 0.61	0.90
PEF (l/sec)	5.25 ± 2.31	5.39 ±2.28	0.96
SVC (I)	3.50 ± 1.17	3.59 ± 1.23	0.91
FEV ₁ /FVC	0.51 ± 0.14	0.51 ± 0.14	0.96
IC (I)	2.65 ± 0.85	2.67 ± 0.87	0.93
Bronchodilator Response (%)++			
FEV ₁	16.3 ± 17.6	16.1 ± 15.2	0.87
FVC	13.2 ± 13.1	11.4 ± 12.0	0.81
Six minute Walk Distance	372.10 ±126.54	394.50 ± 107.91	0.79

Table 2. Baseline and repeat visit intraclass correlation coefficients for objective measures.

Means ± SD

*ICC = Intraclass Correlation

+ COPD = Strata 3&4 only

⁺⁺ Bronchodilator Response calculated as [(post-BD_Pre-BD) / Pre-BD FEV₁] x 100. Analysis was restricted to subjects who had not used any long-acting bronchodilator in the past 48 hours or a LABA within the past 24 hours, Tiotropium within the past 48 hours, or a SABA within 6 hours or ipratropium within the last 8 hours of the baseline and repeat visits (n = 36).

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(59%) had no change in dyspnea, 9 subjects (13%) increased their dyspnea score by greater than 1 and 19 subjects (28%) reported decreased dyspnea of at least 1 grade. Total white blood cell count ICC was 0.76 for all subjects and 0.72 for subjects with COPD supporting the assessment of short-term stability of the subjects between visits.

Measures dependent on patient recall

The presence of CT defined emphysema produced highly reproducible results between visits (Kappa = 0.91) (not shown). The repeatability of self-reported physician diagnosis of emphysema was Kappa = 0.71. Patient recall of physician-diagnosed CB was similar between visits (Kappa = 0.78), but higher than that of CB determined at baseline by the classic definition (Kappa = 0.61). At baseline, 20 subjects (25%) met the classic CB definition, whereas SGRQ identified 41 subjects (48%); agreement between these definitions of CB was relatively poor (Kappa = 0.35). Recall of an asthma diagnosis by a health care professional at baseline and the repeat visit resulted in kappa = 0.57 and Kappa = 0.40 for asthma diagnosed as a child by a health care professional. (Table 3)

Exacerbation frequency is a commonly assessed parameter in COPD. We examined several "definitions" of reported exacerbations including total number of exacerbations of any severity, exacerbations requiring an emergency department (ED) visit or hospitalization, and those

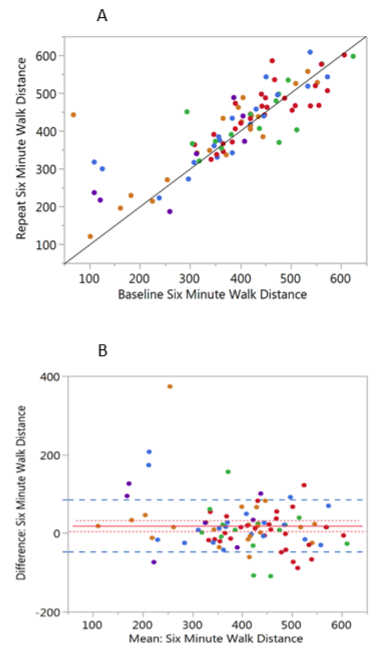


Fig 2. Six minute walk distance. The subjects are color coded by GOLD classification as in Fig 1A) Scatterplot of the six-minute walk distance at the baseline and repeat visits (r = 0.829, p < 0.0001; n = 92). The solid black line is drawn as a line of identity to visualize differences between the baseline and the repeat visits. B) Bland-Altman plot of the mean distance for each subject (baseline and repeat) by the difference between visits. The solid red line = the mean difference, the dotted red line is ± 1 SE and the Dashed blue line is ± 1 SD.

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treated with any medication (corticosteroids or antibiotics). The repeatability of exacerbation recall was modest, but fairly consistent across exacerbation definitions (Table 3). Repeatability increased if the recall of an exacerbation was associated with use of a specific medication such as a corticosteroid or antibiotic. For total exacerbations recall, 48 subjects (71%) remained unchanged, with 59 subjects (87%) unchanged in recall of ED/hospitalization exacerbations.

Variable Name		Kappa Statistics		
	N	Карра	95% CI	
Emphysema HCP Diagnosed	88	0.71	0.55	0.87
CB HCP Diagnosed*	87	0.78	0.61	0.95
CB Classic definition ⁺	95	0.61	0.42	0.80
CB SGRQ**	82	0.61	0.44	0.78
Asthma HCP Diagnosed##	85	0.57	0.36	0.78
Asthma Child Diagnosis	84	0.40	0.10	0.69
Exacerbations in prior 12 months	68	0.42	0.23	0.61
Exacerbations Treated with Any Medications	68	0.45	0.23	0.66
Exacerbations Treated with Corticosteroids	68	0.54	0.31	0.77
Exacerbations Treated with Antibiotics	68	0.58	0.36	0.79
Exacerbations Requiring ED/Hospitalization	68	0.57	0.33	0.82

Table 3. Measures dependent of patient recall: Disease diagnosis and exacerbations.

* CB-Chronic

Bronchitis Diagnosed by a Health Care Provider (HCP)

+ Chronic mucus production and cough for at least 3 months/year for 2 successive years.

** Chronic bronchitis defined as cough and phlegm production several days a week or almost every day from the SGRQ.

Asthma diagnosed as a child by a HCP

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Subjective measures

Subject recall of the age started smoking, average cigarettes smoked per day over the lifetime of smoking and calculated pack-years were evaluated. The baseline and repeat visit values for age started smoking were 16.4 ± 4.5 and 16.0 ± 3.5 respectively (ICC = 0.76), average cigarettes per day 25.4 ± 9.1 and 24.5 ± 9.9 (ICC = 0.79) and calculated pack-years smoked 47.4 ± 19.2 and 49.4 ± 21.6 (ICC = 0.84).

Questionnaires commonly used in COPD showed wide disparity in repeatability (Table 4). Performance ranged from SGRQ-C total score (ICC = 0.94), to MRC dyspnea (kappa = 0.42). The mean change in SGRQ-C between visits was 2.2 units; 34 subjects (55.7%) had no change above the minimum clinically important difference (MCID) of 4 units (Table 5) [14]. The two

Clinical Questionnaire	Baseline	Repeat Visit	ICC* COPD
MRC Dyspnea	1.5 ± 1.1	1.3 ± 1.0	Kappa 0.42 ⁺
Bode Index	2.5 ± 2.2	2.1 ± 1.8	0.84
COPD Assessment Test (CAT)	16.3 ± 8.5	16.2±7.9	0.78
SGRQ-C Total Score	41.8±21.2	40.2 ± 21.5	0.94
FACIT-F Total	111.0 ± 26.1	112.1 ± 27.3	0.91
PSQ Total Score	7.0 ± 4.2	6.9 ± 4.3	0.85
Medical Outcomes SF-12			
SF-12 Gen Health	25.0 ± 11.7	26.5 ± 12.0	0.68
SF-12 Physical Functioning	21.3 ± 12.8	22.7 ± 12.5	0.63

Table 4. ICC values for clinical questionnaires.

Data are Mean ± SD

*ICC = Intraclass Correlation, COPD subjects only

⁺ Kappa Statistic was utilized because of the categorical nature of the data

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Table 5. Change from baseline visit.

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	n	Unchanged ¹	Increased	Decreased
Total Exacerbations	68	48 (71%)	6 (9%)	14 (21%)
Exacerbations requiring ED/Hospitalization	68	59 (87%)	4 (6%)	5 (7%)
SGRQ ²	84	27 (44%)	10 (16%)	24 (39%)
COPD Assessment Test (CAT) ²	65	23 (35%)	21 (32%)	21 (32%)
mMRC ²	98	41 (59%)	9 (13%)	19 (28%)
GOLD PFTs	69	53 (77%)	4 (6%)	12 (17%)
GOLD Combined-mMRC ³	64	44 (69%)	7 (11%)	13 (20%)
GOLD Combined- CAT ³	64	46 (72%)	9 (14%)	9 (14%)

Number of subjects (% of total). n = number changed >MCID: SGRQ = 4; CAT = 2; mMRC = 1. Increase indicates a change from A to B, B to C or C to D

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scores incorporated into the GOLD 2015 and 2017 [15] guidelines (CAT and MRC Dyspnea) appeared to have modest repeatability between visits (ICC = 0.78 and kappa = 0.42, respectively). Change between visits in CAT score ranged from +18 to -11. Among subjects with COPD, 23 of 65 (35.4%) had no change in CAT score (Table 5) above the MCID (2 units). For the MRC dyspnea score, 41 subjects (59.4%) were unchanged (Table 5).

Evaluation of GOLD classification determined solely by PFTs for the COPD subjects demonstrated that 53 of 69 subjects (76.8%) retained their GOLD stage at the repeat visit. (kappa = 0.75). Utilizing the GOLD symptom burden / exacerbation risk criteria [15] with MMRC, 44 of 64 (five subjects had missing mMRC scores) subjects remained unchanged (kappa = 0.54) and with the CAT assessment 46 of 64 (72%) remained the same (kappa = 0.58) (Table 5).

Discussion

This formal, multi-center repeatability study, performed over an average 29-day interim in a representative subset of the entire SPIROMICS cohort, provides an unprecedented assessment of the variability of the instruments used to characterize COPD subjects. Repeatability of pulmonary function testing was strongest (ICC >0.90) and subject recall of a childhood diagnosis of asthma weakest (kappa = 0.40), consistent with a trend of greater repeatability of objective measures relative to those dependent on participant recall or subjectivity.

Our analytic plan was designed specifically to improve estimation of the reliability of clinically relevant predictors of COPD severity and activity. Inherent variability in a predictor, due either to measurement error or to short-term biological fluctuations, may bias estimates of the association between the predictor and an outcome [16]. Estimation of the reliability of a predictor, such as the intraclass correlation, allows correction for the bias using regression calibration. This is important in defining COPD subsets for analysis and in evaluating disease progression.

These results relate to the complexity of what we call COPD. Historically, the disease has been conceptualized as having a slow, often variably progressive decline in lung function. [17–19]. However, recent data suggests that the slope of decline in FEV₁% predicted slows in advanced disease[20] and that correlations between lung function and other measures such as exacerbations and symptoms are not always strong [21, 22]. Nevertheless, FEV₁ remains the objective gold standard for assessing disease severity, progression, and treatment efficacy. Our finding that all PFTs had an ICC >0.90 supports the use of PFTs as a primary outcome variable. These data are consistent with other reports demonstrating the high repeatability of spirometric values between visits of short intervals over a range of disease severities [12, 23]. Our

mean change in FEV₁ (20.4 mL overall, 30.7 mL for men, 1.9 mL for women), compares favorably with results from the Lung Health Study (LHS) at screening visits 21 days apart, which reported a coefficient of variation of 4% (changes of 14.3 mL for men, 4.5 mL for women)[23]. Similar results were reported for FEV₁ in the National Emphysema Treatment Trial (NETT) for PFTs conducted within 60 days of each other [12]. We extend those finding by showing that IC, an important driver of exercise limitation, was as repeatable as other spirometric parameters (ICC = 0.93) [24–26]. IC decreases during exercise in COPD, and is responsive to bronchodilator therapy [27, 28]. Bronchodilator response is influenced by many factors including pre-bronchodilator FEV₁ and actual withholding of confounding bronchodilator medications [29]. In our study, withholding bronchodilators was not required, but extensive questions were asked on drug use and time of administration. Our reported bronchodilator response was calculated in subjects who withheld bronchodilators (65 out of 98 subjects) for sufficient time before PFT testing, potentially representing a more accurate picture of bronchodilator responsiveness.

Fully assessing the full clinical picture of COPD requires additional factors and composite scores dependent on patient recall and subjective evaluation [30]. We found that most of the validated questionnaires had lower ICCs than the PFTs. Among them, the most repeatable was the SGRQ-C, which is tailored for use in chronic airflow limitation and is responsive to changes in disease activity [31, 32]. Repeatability of SGRQ-C has been reported as ICC = 0.92 in 40 subjects evaluated at a 2 week interval [31], in good agreement with our results. Each questionnaire has a different specified recall time period, ranging from none specified (SGRQ-C) to 7 days for the FACIT-F and current assessment for the CAT, which could affect repeatability. Nevertheless, we found significant correlations between the utilized questionnaire scores (data not shown), suggesting that they are measuring similar aspects of the disease. This relationship is not surprising because all aim to assess the functional impact of COPD on quality of life. Our finding that repeatability of subject recall of age started smoking was similar to assessing average number of cigarettes smoked over their life-time, is consistent with an extensive internet based survey of tobacco exposure and risk [33]. They found age first started smoking cigarettes slightly more consistent (ICC = 0.85) than calculated pack-years (ICC = 0.76). Despite our finding of a few subjects with very different recall of pack-years smoked, overall, pack-years seems relatively reliable, but of unknown accuracy.

Finally, these results provide insights into the use of patient recall of exacerbation history to predict susceptibility to future exacerbations. Our results on short-term repeatability of recalling COPD exacerbations before enrollment (total exacerbations, kappa = 0.42), was somewhat higher when defined by treatment with corticosteroids or antibiotics (kappa = 0.54 and 0.58, respectively). These data are similar to reported concordance of cardiovascular events, in that the more defined the event, the greater the reliability [34]. It was surprising that recall of exacerbations requiring an emergency department visit or hospitalization was not significantly higher. The frequency of exacerbations, and not the severity, may be the most important factor in patient recall inaccuracies [35]. In the ECLIPSE study, patient recall of the number of exacerbations in the year before enrollment was the single strongest predictor of future exacerbations [22] highlighting the importance of exacerbation recall for COPD studies. However, our current results imply that using patient recall of exacerbation frequency as an enrollment criterion for short-term clinical trials may lead to substantial variability in outcomes. The inaccuracy of patient recall of exacerbation rate was recently highlighted when compared with singlephysician chart review or a central adjudication committee [35]. It is surprising that recall of hospitalization or ED visit was not consistent, highlighting the need for exacerbation documentation in determining GOLD status and associated treatment choices.

A key feature of our analysis is its implications for the GOLD combined assessment of COPD, which is employed both for assessment of disease severity and for treatment recommendations. We found a higher reliability (as judged by ICC) for the GOLD classification using PFTs. Incorporation of symptom scores (MMRC and CAT) into GOLD classification for symptoms and exacerbation risk increases variability between visits. Using the 2015 GOLD combined assessment criteria, 31% of the subjects changed by at least one GOLD level using GOLD-MMRC and 18% changed using GOLD-CAT. We recognize that there are two possible interpretations of the more limited repeatability of symptom-based scores compared to PFTs. It is possible that symptom scores are more sensitive than PFTs for clinical changes and provide a more sensitive indicator of current illness. This is reflected in the updated GOLD criteria [15, 36]. Symptom scores also reflect an integrated assessment of a multi-organ disease and comorbidities that are common in these subjects. However, the observation that parameters dependent upon subject recall also have poorer consistency upon repeat evaluation suggests that a component of the variation in symptoms, in the absence of changes in spirometry, may be a reflection of the subjective nature of the assessment. Further work is required to best delineate the precision of change in symptom scores as endpoints for clinical trials in COPD.

As with all studies, there are limitations. This Repeatability Substudy evaluated differences in participant response over an interval of 28.9 ± 9.4 days (mean \pm SD). We did not adjust for variation in time between visits. It is also unknown how a longer or shorter time would correlate with the results reported here. Though demographics of the Substudy were consistent with the total SPIROMICS cohort, only 7 of the 11 clinical centers contributed data to this Substudy, so these results may not reflect repeatability across all of the centers. Also, we cannot rule out an effect of training [37], as some measures improved at the repeat visit. Perhaps most crucially, these subjects may not be representative of the general COPD population.

In summary, we demonstrate that in the SPIROMICS Repeatability Substudy, the repeatability of supervised objective measures was strongest. PFTs demonstrated the highest ICC values and recall of exacerbations had some of the lowest kappa statistics. Reliability of questionnaires was consistent with literature reports. However, significant numbers of subjects had variation between visits above recommended MCIDs, which notably affected GOLD staging for symptoms and exacerbation risk incorporating either the CAT or MMRC. Within-subject variability must be accounted for in interpreting phenotype assignments or disease progression. We feel that these data will aid the design and interpretation of longitudinal COPD studies. Examining multiple types of parameters widely used in COPD assessment should also help in the development of clinical practice guidelines.

Supporting information

S1 Fig. PFTs and bland-altman plots. Subjects (n = 96) are color coded by GOLD stratification (using PFT values only). GOLD 0 = red, GOLD 1 = green, GOLD 2 = Blue, GOLD 3 = orange and GOLD 4 = Purple. The solid red line = the mean difference between the base-line and repeat visit values, the dotted red line is \pm 1 SE and the Dashed blue line is \pm 1 SD. A) Post-bronchodilator FVC, B) Post-bronchodilator FVC Bland-Altman Plot, C) Post-bronchodilator FEV₁/FVC, D) Post-bronchodilator FEV₁/FVC Bland-Altman Plot, E) Post-bronchodilator Inspiratory Capacity and F) Post-bronchodilator Inspiratory Capacity Bland-Altman Plot.

(TIF)

S1 Study Information. (DOCX)

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References

- Couper D, LaVange LM, Han M, Barr RG, Bleecker E, Hoffman EA, et al. Design of the Subpopulations and Intermediate Outcomes in COPD Study (SPIROMICS). Thorax. 2014 May; 69(5):491–4. https:// doi.org/10.1136/thoraxjnl-2013-203897 PMID: 24029743. Pubmed Central PMCID: PMC3954445. Epub 2013/09/14. eng.
- Bui MK-L, Nyberg DA, Maltais DF, Saey DD. Functional Tests in Chronic Obstructive Pulmonary Disease Part 2: Measurement Properties. Ann Am Thorac Soc. 2017 Feb 28. https://doi.org/10.1513/AnnalsATS.201609-734AS PMID: 28244801.
- 3. Anderson WH, O'Neal WK, Doerschuk CM, Carretta EE, Couper DJ, Tashkin D, et al. Short-Term Stability Of Pulmonary Function And Clinical Measures In COPD Using A Cohort From SPIROMICS

(Subpopulations And Intermediate Outcome Measures In COPD Study) [abstract]. 193: Am J Respir Crit Care Med; 2016. p. A3515.

- Jones PW, Harding G, Berry P, Wiklund I, Chen WH, Kline Leidy N. Development and first validation of the COPD Assessment Test. The European respiratory journal. 2009 Sep; 34(3):648–54. https://doi. org/10.1183/09031936.00102509 PMID: 19720809. Epub 2009/09/02. eng.
- Meguro M, Barley EA, Spencer S, Jones PW. Development and Validation of an Improved, COPD-Specific Version of the St. George Respiratory Questionnaire. Chest. 2007 Aug; 132(2):456–63. <u>https://doi.org/10.1378/chest.06-0702</u> PMID: 17646240. Epub 2007/07/25. eng.
- Al-shair K, Muellerova H, Yorke J, Rennard SI, Wouters EF, Hanania NA, et al. Examining fatigue in COPD: development, validity and reliability of a modified version of FACIT-F scale. Health and quality of life outcomes. 2012; 10:100. https://doi.org/10.1186/1477-7525-10-100 PMID: 22913289. Pubmed Central PMCID: PMC3491053. Epub 2012/08/24. eng.
- Buysse DJ, Reynolds CF 3rd, Monk TH, Berman SR, Kupfer DJ. The Pittsburgh Sleep Quality Index: a new instrument for psychiatric practice and research. Psychiatry research. 1989 May; 28(2):193–213. PMID: 2748771. Epub 1989/05/01. eng.
- Ware J Jr., Kosinski M, Keller SD. A 12-Item Short-Form Health Survey: construction of scales and preliminary tests of reliability and validity. Medical care. 1996 Mar; 34(3):220–33. PMID: 8628042. Epub 1996/03/01. eng.
- 9. Ferris BG. Epidemiology Standardization Project (American Thoracic Society). The American review of respiratory disease. 1978 Dec; 118(6 Pt 2):1–120. PMID: 742764. Epub 1978/12/01. eng.
- Kim V, Crapo J, Zhao H, Jones PW, Silverman EK, Comellas A, et al. Comparison between an alternative and the classic definition of chronic bronchitis in COPDGene. Annals of the American Thoracic Society. 2015 Mar; 12(3):332–9. https://doi.org/10.1513/AnnalsATS.201411-518OC PMID: 25575351. Pubmed Central PMCID: PMC4418311. Epub 2015/01/13. eng.
- Vestbo J, Hurd SS, Agustí AG, Jones PW, Vogelmeier C, Anzueto A, et al. Global Strategy for the Diagnosis, Management, and Prevention of Chronic Obstructive Pulmonary Disease. Am J Respir Crit Care Med. 2013; 187(4):347–65. https://doi.org/10.1164/rccm.201204-0596PP PMID: 22878278.
- Herpel LB, Kanner RE, Lee SM, Fessler HE, Sciurba FC, Connett JE, et al. Variability of spirometry in chronic obstructive pulmonary disease: results from two clinical trials. American journal of respiratory and critical care medicine. 2006 May 15; 173(10):1106–13. https://doi.org/10.1164/rccm.200506-975OC PMID: 16497996. Pubmed Central PMCID: PMC2662942. Epub 2006/02/25. eng.
- **13.** Puhan MA, Chandra D, Mosenifar Z, Ries A, Make B, Hansel NN, et al. The minimal important difference of exercise tests in severe COPD. The European respiratory journal. 2011 Apr; 37(4):784–90. https://doi.org/10.1183/09031936.00063810 PMID: 20693247. Epub 2010/08/10. eng.
- Jones PW. St. George's Respiratory Questionnaire: MCID. Copd. 2005 Mar; 2(1):75–9. PMID: 17136966. Epub 2006/12/02. eng.
- Vogelmeier CF, Criner GJ, Martinez FJ, Anzueto A, Barnes PJ, Bourbeau J, et al. Global Strategy for the Diagnosis, Management, and Prevention of Chronic Obstructive Lung Disease 2017 Report. GOLD Executive Summary. American journal of respiratory and critical care medicine. 2017 Mar 01; 195 (5):557–82. https://doi.org/10.1164/rccm.201701-0218PP PMID: 28128970. Epub 2017/01/28. eng.
- Rosner B, Willett WC, Spiegelman D. Correction of logistic regression relative risk estimates and confidence intervals for systematic within-person measurement error. Statistics in medicine. 1989 Sep; 8 (9):1051–69; discussion 71–3. PMID: 2799131. Epub 1989/09/01. eng.
- Rennard SI, Vestbo J. Natural histories of chronic obstructive pulmonary disease. Proceedings of the American Thoracic Society. 2008 Dec 15; 5(9):878–83. https://doi.org/10.1513/pats.200804-035QC PMID: 19056710. Pubmed Central PMCID: PMC2720106. Epub 2008/12/06. eng.
- Lange P, Halpin DM, O'Donnell DE, MacNee W. Diagnosis, assessment, and phenotyping of COPD: beyond FEV1. International journal of chronic obstructive pulmonary disease. 2016; 11 Spec Iss:3–12. https://doi.org/10.2147/COPD.S85976 PMID: 26937185. Pubmed Central PMCID: PMC4765947. Epub 2016/03/05. eng.
- Fletcher C, Peto R. The natural history of chronic airflow obstruction. BMJ. 1977 1977-06-25 07:00:00; 1(6077):1645–8. PMID: 871704
- Tantucci C, Modina D. Lung function decline in COPD. International journal of chronic obstructive pulmonary disease. 2012; 7:95–9. https://doi.org/10.2147/COPD.S27480 PMID: 22371650. Pubmed Central PMCID: PMC3282601. Epub 2012/03/01. eng.
- Curtis JR, Deyo RA, Hudson LD. Pulmonary rehabilitation in chronic respiratory insufficiency. 7. Healthrelated quality of life among patients with chronic obstructive pulmonary disease. Thorax. 1994 Feb; 49 (2):162–70. PMID: 8128407. Pubmed Central PMCID: PMC474337. Epub 1994/02/01. eng.

- Hurst JR, Vestbo J, Anzueto A, Locantore N, Mullerova H, Tal-Singer R, et al. Susceptibility to exacerbation in chronic obstructive pulmonary disease. N Engl J Med. 2010 Sep 16; 363(12):1128–38. https://doi.org/10.1056/NEJMoa0909883 PMID: 20843247. Epub 2010/09/17. eng.
- Enright PL, Beck KC, Sherrill DL. Repeatability of spirometry in 18,000 adult patients. American journal of respiratory and critical care medicine. 2004 Jan 15; 169(2):235–8. https://doi.org/10.1164/rccm. 200204-347OC PMID: 14604836. Epub 2003/11/08. eng.
- Ramon MA, Ferrer J, Gimeno-Santos E, Donaire-Gonzalez D, Rodriguez E, Balcells E, et al. Inspiratory capacity-to-total lung capacity ratio and dyspnoea predict exercise capacity decline in COPD. Respirology (Carlton, Vic). 2016 Apr; 21(3):476–82. https://doi.org/10.1111/resp.12723 PMID: 26714424. Epub 2015/12/31. eng.
- Klooster K, ten Hacken NH, Hartman JE, Sciurba FC, Kerstjens HA, Slebos DJ. Determining the Role of Dynamic Hyperinflation in Patients with Severe Chronic Obstructive Pulmonary Disease. Respiration; international review of thoracic diseases. 2015; 90(4):306–13. https://doi.org/10.1159/000439056 PMID: 26352833. Epub 2015/09/10. eng.
- Zhang Y, Sun XG, Yang WL, Tan XY, Liu JM. Inspiratory fraction correlates with exercise capacity in patients with stable moderate to severe COPD. Respiratory care. 2013 Nov; 58(11):1923–30. https:// doi.org/10.4187/respcare.01927 PMID: 23592787. Epub 2013/04/18. eng.
- Yan S, Kaminski D, Sliwinski P. Reliability of inspiratory capacity for estimating end-expiratory lung volume changes during exercise in patients with chronic obstructive pulmonary disease. American journal of respiratory and critical care medicine. 1997 Jul; 156(1):55–9. https://doi.org/10.1164/ajrccm.156.1. 9608113 PMID: 9230726. Epub 1997/07/01. eng.
- Celli BR, Decramer M, Lystig T, Kesten S, Tashkin DP. Longitudinal inspiratory capacity changes in chronic obstructive pulmonary disease. Respiratory research. 2012; 13:66. https://doi.org/10.1186/ 1465-9921-13-66 PMID: 22866681. Pubmed Central PMCID: PMC3443002. Epub 2012/08/08. eng.
- Calverley PM, Albert P, Walker PP. Bronchodilator reversibility in chronic obstructive pulmonary disease: use and limitations. The Lancet Respiratory medicine. 2013 Sep; 1(7):564–73. https://doi.org/10.1016/S2213-2600(13)70086-9 PMID: 24461617. Epub 2014/01/28. eng.
- Salzman SH. Which pulmonary function tests best differentiate between COPD phenotypes? Respiratory care. 2012 Jan; 57(1):50–7; discussion 8–60. <u>https://doi.org/10.4187/respcare.01585</u> PMID: 22222125. Epub 2012/01/10. eng.
- Jones PW, Quirk FH, Baveystock CM, Littlejohns P. A self-complete measure of health status for chronic airflow limitation. The St. George's Respiratory Questionnaire. The American review of respiratory disease. 1992 Jun; 145(6):1321–7. <u>https://doi.org/10.1164/ajrccm/145.6.1321</u> PMID: <u>1595997</u>. Epub 1992/06/01. eng.
- 32. Puhan MA, Guyatt GH, Goldstein R, Mador J, McKim D, Stahl E, et al. Relative responsiveness of the Chronic Respiratory Questionnaire, St. Georges Respiratory Questionnaire and four other healthrelated quality of life instruments for patients with chronic lung disease. Respiratory medicine. 2007 Feb; 101(2):308–16. <u>https://doi.org/10.1016/j.rmed.2006.04.023</u> PMID: <u>16782320</u>. Epub 2006/06/20. eng.
- Brigham J, Lessov-Schlaggar CN, Javitz HS, Krasnow RE, McElroy M, Swan GE. Test-retest reliability of web-based retrospective self-report of tobacco exposure and risk. Journal of medical Internet research. 2009; 11(3):e35. https://doi.org/10.2196/jmir.1248 PMID: 19674962. Pubmed Central PMCID: PMC2762856. Epub 2009/08/14. eng.
- Heckbert SR, Kooperberg C, Safford MM, Psaty BM, Hsia J, McTiernan A, et al. Comparison of selfreport, hospital discharge codes, and adjudication of cardiovascular events in the Women's Health Initiative. American journal of epidemiology. 2004 Dec 15; 160(12):1152–8. https://doi.org/10.1093/aje/ PMID: 15583367. Epub 2004/12/08. Eng.
- 35. Frei A, Siebeling L, Wolters C, Held L, Muggensturm P, Strassmann A, et al. The Inaccuracy of Patient Recall for COPD Exacerbation Rate Estimation and its Implications: Results from Central Adjudication. Chest. 2016 Jul 8; 150(4):860–8. https://doi.org/10.1016/j.chest.2016.06.031 PMID: 27400907. Epub 2016/07/13. Eng.
- Vogelmeier C. Global Strategy for the Diagnosis, Management and Prevention of COPD, Global Initiative for Chronic Obstructive Lung Disease (GOLD) 2015. 2015.
- Knox AJ, Morrison JF, Muers MF. Reproducibility of walking test results in chronic obstructive airways disease. Thorax. 1988 May; 43(5):388–92. PMID: <u>3194867</u>. Pubmed Central PMCID: PMC461257. Epub 1988/05/01. eng.