Implications of using the GLI-2012, GOLD and Australian COPD-X recommendations in assessing the severity of airflow limitation on spirometry among an Indigenous population with COPD: an Indigenous Australians perspective study

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ABSTRACT

Background Assessment of airflow limitation (AFL) is crucial in the clinical evaluation of patients with chronic obstructive pulmonary disease (COPD). However, in the absence of normative reference values among adult Australian Indigenous population, the implications of utilising the Global Lung Function Initiative (GLI-2012), Global Initiative for Chronic Obstructive Lung Disease (GOLD) and the Australian concise COPD-X recommended severity classifications is not known. Moreover, spirometry values (forced vital capacity (FVC) and forced expiratory volume in 1 s (FEV1)) are observed to be 20%–30% lower in an apparently healthy Indigenous population in comparison to Caucasian counterparts.

Methods Adult Indigenous patients diagnosed to have COPD on spirometry (postbronchodilator (BD) FEV1/FVC <0.7 ((GOLD, (COPD-X)) and ≤lower limit of normal (others/mixed reference equations) for GLI-2012) were assessed for AFL severity classifications on Post-BD FEV1 values (mild, moderate, severe, very severe) as per the recommended classifications.

Results From a total of 742 unique patient records of Indigenous Australians, 253 were identified to have COPD via GOLD/COPD-X criteria (n=238) or GLI-2012 criteria (n=238) with significant agreement between criteria (96%, κ=0.901). Of these, the majority were classified as having moderate or severe/very-severe AFL with significant variability across classification criteria (COPD-X (40%–43%), GOLD (33%–65%), GLI-2012 (18%–75%)). The FVC and FEV1 values also varied significantly between classification criterion (COPD-X/GOLD/GLI-2012) within the same AFL category, with COPD-X ‘moderate’ AFL almost matching ‘severe’ AFL categorisation by GOLD or GLI-2012.

Conclusions Health professionals caring for Indigenous patients with COPD should be aware of the clinical implications and consequences of utilising various recommended AFL classifications in the absence of validated spirometry reference norms among adult Indigenous patients.

INTRODUCTION

Chronic obstructive pulmonary disease (COPD) is one of the leading causes of morbidity and mortality among the Aboriginal and Torres Straits Islander populations of Australia (henceforth respectfully referred to as Indigenous Australians or Indigenous people or patients), with a much higher prevalence reported in the Northern Territory (NT) Indigenous population. Moreover, hospital admission rates are also reported to be significantly higher for the adult Indigenous patients and COPD is one of the potentially preventable causes of admissions.
in this population. Emerging evidence in the literature suggests that Indigenous Australians not only have a higher prevalence of COPD, but also tend to exhibit other concurrent pulmonary conditions, in particular the presence of bronchiectasis, along with significantly lower lung function parameters (LFPs) in comparison to their non-Indigenous counterparts.

In the clinical assessment of patients presenting with symptoms of COPD, the lung function tests/spirometry play an integral role in the diagnosis, assessment of severity of airflow limitation (AFL) and in guiding therapeutic interventions. However, lung function norms are yet to be established for the adult Indigenous population. The limited available evidence in the literature suggests that Indigenous Australians may have up to 20%–30% lower values for forced vital capacity (FVC) and forced expiratory volume in 1 s (FEV₁) in comparison to Australian Caucasians. Furthermore, although the Global Lung Function Initiative (GLI-2012) spirometric reference norms are shown to fit various ethnic groups including the general adult Australian population, due to lack of Indigenous Australians representation in the GLI-2012 contemporary adult Australian data, it is unclear if the GLI-2012 data fit the adult Indigenous Australian population. Recently, smaller studies have demonstrated that the GLI-2012 norms may not be appropriate for the Australian Indigenous adults regardless of which ethnicity option is selected, including ‘others/mixed’. Yet, the current recommendations are to use GLI-2012 ‘others/mixed’ equations among the adult Indigenous Australians, Torres Strait Islanders, New Zealand Māori and Pacific Island populations.

Several well-established recommendations, including the global strategy for the diagnosis, management, and prevention of COPD (global initiative for chronic obstructive lung disease (GOLD)), have been developed for the purpose of classifying the severity of AFL among patients with COPD (GOLD 1 to 4) by spirometry in the overall holistic management. In the Australian context, the concise tool for COPD-X is the recommended/endorsed tool that is widely adopted in assessing the severity of AFL by spirometry in order to facilitate in the stepwise management of patients with COPD. The COPD-X guidelines have also been endorsed by the Royal Australian College of General Practitioners for use at the primary healthcare level.

Nevertheless, the implications of utilising these recommended spirometry parameters in assessing/classifying the severity of AFL among Indigenous Australians have not been investigated. Moreover, the spirometry cut-off values to assess the severity of AFL differ between the two recommendations (ie, ‘GOLD’ vs ‘COPD-X’). Therefore, the adoption of either of these tools in assessing the severity of AFL among Indigenous patients with COPD may have differing implications due to the following three reasons. First, Indigenous Australian adults appear to have up to 20%–30% lower values for FEV₁ in comparison to Australian Caucasians. Second, there are no well-established spirometry norms for the adult Indigenous population, and thus reference is made against the Caucasian population. Finally, there is a higher prevalence of COPD and COPD/Bronchiectasis among Indigenous Australians. Taking into consideration these aforementioned factors, it is inevitable that the severity of COPD, in terms of AFL, will be either under or overestimated, or potentially lead to misclassification of COPD stages, which in turn may have significant implications on the short and long term management of COPD among Indigenous patients.

Hence, it may be important to evaluate the relevance/implications of applying the GLI-2012, GOLD and COPD-X recommendations to an Indigenous Australian population and assess how the spirometry parameters (namely FEV₁ and FEV₁/FVC) may display among patients with a clinical diagnosis of COPD. Therefore, this study endeavours to investigate the relevance and implications of AFL severity classifications in an Australian Indigenous population diagnosed to have COPD in the Top End Health Service (TEHS) region of the NT of Australia.

POPULATION AND METHODS

Setting and ethics

This study was conducted at the Respiratory and Sleep service based at the Royal Darwin hospital/Darwin respiratory and sleep health/Darwin private hospital in the TEHS region of the NT of Australia. In the NT, approximately 30% of the population self-identify as Indigenous Australian, the highest proportion compared with all other Australian states and territories.

The study participants included in this study are inclusive of the data sets of participants from our previous reports assessing factors influencing and implications of chest radiological findings and LFPs in the adult Indigenous Australians residing in the Top End NT of Australia. This study was conducted and reported according to strengthening reporting of health research involving Indigenous people.

Patient and public involvement

As the study was retrospective in nature, patients’ participation or individual consent from the study participants was not required. In the public involvement perspective, local institute Indigenous Australian representatives were consulted in reviewing and approving this manuscript for its design, conduct, reporting, dissemination of our research work, in particular, the appropriateness and respect in relation to the Indigenous context represented in this study.

Study participants and inclusion criteria

The study participants included were Indigenous Australian patients residing in the TEHS region of the NT of Australia who had undergone spirometry testing.
between 2012 and 2020. Patients were referred for spirometry by primary health practitioners, respiratory specialist and other specialist physicians as a part of routine clinical care. A total of 1350 lung function tests were recorded by Indigenous patients during the study period, of which 965 met test acceptability criteria. This assessment only used the earliest recorded and acceptable for session quality lung function test of Indigenous patients, resulting in a total 742 tests available for analysis.

**Lung function testing**

All spirometry testing were performed according to the American thoracic and the European respiratory societies guidelines/recommendations, including calibration of equipment and quality assurance. The tests were performed via a portable single-breath diffusing capacity device ‘EasyOne Pro, ndd Medical Technologies’. All volume-time and flow-volume graphs were individually assessed for acceptability for session quality. Only spirometric tests that were graded as acceptable for session quality were included in this study. More details are available from previous reports from our centre.

When appropriate, all patients undergoing elective spirometry testing were instructed to refrain from smoking for at least 2–4 hours prior to spirometry testing and to avoid using airway directed inhaled pharmacotherapy during the preceding 6–12 hours of spirometry testing, and among patients taking some forms of longer acting inhaled pharmacotherapy to avoid for 24 hours prior to testing. According to the GOLD criteria used in earlier burden of obstructive lung disease studies, bronchodilator (BD) responsiveness for spirometry parameters were assessed 15–20 min after inhalation of 400 µg of salbutamol via a spacer. The presence of a clinically significant BD responsiveness was determined if there was a post-BD increase in FEV1 of ≥12% and ≥200 mL from the pre-BD FEV1. For the purpose of this study, only post-BD percent predicted values for FEV1 and FEV1/FVC ratio values were reported. In our centre, faced with the lack of specific spirometric norms for the adult Australian Indigenous population, the predicted values were calculated using the third National health and nutrition examination survey (NHANES) Caucasian reference sets (‘other ethnicity’). For comparison with GLI-2012, the dataset was run through the GLI-2012 dataset conversion app using ‘others/mixed’ reference ethnicity, with resultant lower limits of normal (LLN) and z-scores used throughout this study. If the study participants were identified to have undergone multiple LFTs during the study period, the earliest and acceptable test for session quality was utilised for analysis.

**COPD diagnosis**

To ascertain the presence of COPD, post-BD FEV1/FVC <0.7 was used for GOLD and COPD-X, and ≤LLN was used for GLI-2012.

**COPD AFL severity assessment criteria**

Table 1 illustrates the COPD severity criteria’s according to three international recommendations. In brief, for GLI-2012 four grading AFL severity (based on post-BD FEV1 z-scores) were used. For GOLD, four grading AFL severity (based on percentage predicted post-BD FEV1 values) were used. For COPD-X, three grading AFL severity (based on the percentage predicted post-BD FEV1 values) were used. Both GOLD and COPD-X used the NHANES Caucasian reference set for ‘other ethnicity’ in line with standard practice.

**Clinical data collection**

As per standard protocol, patients’ age (on the date of testing), sex, height, weight and body mass index (BMI, kg/m2) were recorded. Smoking history, indication for spirometry and the patients’ electronic medical records were reviewed to document the clinical diagnosis of

| Table 1 COPD airflow limitation severity criteria’s according to three recommendations |
|-----------------------------------------------|-----------------|----------------|
| **Recommended tools** | **Severity categories** | Post-BD FEV1/FVC (absolute) | Post-BD FEV1 |
|------------------------|------------------------|-----------------|--------------|
| **GOLD** | Mild | <0.7 | ≥80% |
| | Moderate | <0.7 | 50%–79% |
| | Severe | <0.7 | 30%–79% |
| | Very severe | <0.7 | <30% |
| **COPD-X** | Mild | <0.7 | 60%–79% |
| | Moderate | <0.7 | 40%–59% |
| | Severe | <0.7 | <40% |
| **GLI** | Mild | ≤LLN | z-score ≥ −2 |
| | Moderate | ≤LLN | −2.0≥−3.0 |
| | Severe | ≤LLN | −3.0≥−4.0 |
| | Very severe | ≤LLN | z-scores ≤−4.0 |

BD, bronchodilator; COPD, chronic obstructive pulmonary disease; FEV1, forced expiratory volume in 1 s; FVC, forced vital capacity; GLI, global lung function imitative; GOLD, global initiative for chronic obstructive lung disease; LLN, lower limit of normal.
COPD. The patient was noted as deceased if it was recorded in the medical records and no linkage to death register was undertaken.

**Statistical analysis**
Continuous parameters were tested for normality via the Shapiro-Wilks distribution test, with BMI and smoking pack-years the only parameters displaying a non-parametric distribution. Non-parametric parameters were presented as medians (IQR), normally distributed parameters as means (95% CIs) and categorical parameters as numbers (%). Clinical characteristics were stratified by severity category and compared across classification guidelines (COPD-X vs GOLD vs GLI) using univariate linear regression for normally distributed parameters, quantile regression for non-parametric parameters and χ² test for categorical parameters. Cohen’s Kappa (κ) was used to determine agreement in COPD classification between GOLD/COPD-X methods and GLI-2012, and between these criteria and what was reported in the patients’ medical records. LFPs were stratified by severity category and tested for equality across recommended classification via univariate linear regression. All data were analysed in STATA IC V.15 (StataCorp) and alpha was set to 0.05 throughout.

**RESULTS**

**Clinical data**
A total of 742 unique patient records were available for analysis, within which a total of 253 patients were identified to have COPD on spirometry, with significant agreement between the two methods used (FEV₁/FVC<0.7 (n=238) or FEV₁/FVC ≤LLN (n=238), κ=0.907, agreement 96%). Agreement between spirometry criterion and the clinical diagnosis of COPD as recorded in the medical records was significantly lower (GOLD/COPD-X and medical record, κ=0.44, agreement 74% and GLI-2012 and medical record, κ=0.407, agreement 73%). The majority of patients (51%) were male with a mean age 53.6 years (95% CI 52.1 to 55.2), median BMI of 23.04 kg/m² (IQR 19.43–27.96) and a high prevalence of either current (51%) or former (40%) smokers with no significant differences between those identified with COPD via GOLD/COPD-X or GLI-2012 (table 2).

**Spirometry results**
Of the patients with COPD identified, the proportion categorised as severe/very severe was significantly higher using the GLI-2012 criteria (table 3). Patients in the GLI-2012 ‘very severe’ category were significantly younger than patients in the ‘severe’ or ‘very severe’ categories according to COPD-X and GOLD (table 4). Distribution of BMI and smoking status did not differ across AFL severity categories. Although overall the prevalence of deceased patients was similar between COPD-X/GOLD/GLI-2012, (16% (n=38), 16% (n=38), and 15% (n=35), respectively), the distribution of mortality across AFL severity categories significantly differed. Of the 38 deceased in COPD-X/GOLD classification, 31% were in the mild or moderate category according to COPD-X and 16% according to GOLD, whereas of the 35 deceased according to GLI-2012 classification, 12% were in the mild or moderate severity categories (table 3).

| Variables          | Clinical parameters | COPD-X/GOLD | GLI-2012 |
|--------------------|---------------------|-------------|----------|
|                    |                     | n=238       | n=238    |
| Sex                | Male                | 127 (53%)   | 123 (52%)|
| Age                | Years (mean (95% CI)) | 54.67 (53.13 to 56.2) | 52.69 (51.15 to 54.23) |
| Corpulence and status | BMI (median (IQR)) | 22.86 (19.38–27.89) | 22.91 (19.04–27.89) |
|                    | Underweight (BMI <18.5 kg/m²) | 48 (20%) | 51 (22%) |
|                    | Normal weight (BMI 18.5–24.9 kg/m²) | 99 (42%) | 95 (40%) |
|                    | Overweight (BMI 25–29.9 kg/m²) | 50 (21%) | 49 (21%) |
|                    | Obese (BMI ≥30 kg/m²) | 40 (17%) | 42 (18%) |
| Smoking status     | Never smoker        | 20 (8%)     | 17 (7%)  |
|                    | Former smoker       | 95 (40%)    | 98 (41%) |
|                    | Current smoker      | 123 (52%)   | 123 (52%)|
|                    | Smoking pack years (median (IQR)) | 21.5 (7–40) | 20 (6.13–40) |
| Mortality          | Deceased (all cause) | 38 (16%)    | 35 (15%) |
|                    | Deceased age (years (mean (95% CI)) | 57.79 (53.97 to 61.61) | 57.15 (53.08 to 61.22) |
| COPD               | Clinical history of COPD | 161 (70%)   | 156 (68%) |

Fifteen patients were identified via FEV₁/FVC <0.7 who were not identified via FEV₁/FVC <LLN, and vice versa 15 patients were identified via FEV₁/FVC >LLN and not by FEV₁/FVC <0.7.
BMI, body mass index; COPD, chronic obstructive pulmonary disease; FEV₁, forced expiratory volume in 1 s; FVC, forced vital capacity; GLI, global lung function imitative; GOLD, global initiative for chronic obstructive lung disease; LLN, lower-limit of normal.
 Spirometry parameters according to different recommended classifications

Spirometry parameters differed significantly within severity categories across the three recommended classifications (table 5, figure 1). Z-scores of post-BD FVC and FEV₁ were on average one unit lower using COPD-X recommendation for each of the mild, moderate and severe categories compared with both GOLD and GLI-2012 categorisation and for FEV₁/FVC in the severe category. Absolute and percentage predicted values for LFPs were significantly lower across all severity categories when using the COPD-X classification. GOLD and GLI-2012 classifications showed significant overlap of mean LFPs in both the mild and moderate categories; however, in the severe and very severe categories GOLD values were significantly lower. A separate analysis of moderate COPD-X against severe GLI-2012 and GOLD showed that for the majority of LFPs moderate COPD-X did not significantly differ from severe GLI-2012 classification (online supplemental appendix 1).

DISCUSSION

To the best of the authors’ knowledge, this is the first study to demonstrate the clinical implications of classifying severity of AFL by adopting various spirometry COPD severity criterion in an Indigenous population with COPD, especially from the NT of Australia. This study has highlighted that irrespective of which of the three recommended AFL classifications used (COPD-X, GOLD, GLI) (<0.7, ≤LLN), most Indigenous patients with COPD who are referred for lung function testing in a specialist centre, will likely fall into either the ‘severe’ or ‘very severe’ category. Moreover, the FVC and FEV₁ values vary significantly between the three recommended severity classification criteria’s.

Spirometry parameters are crucial in day-to-day clinical practice, not only for the accurate diagnosis and classifying the severity of AFL, but also in the management of COPD with airway directed inhaled pharmacotherapy. While it is possible the high prevalence of ‘severe’ AFL categorisation among Indigenous Australian patients with COPD accurately represents the severity of the underlying disease as noted in this study, it is also plausible that these high rates of ‘severe’ categorisation are an artefact of the use of reference values drawn from non-Indigenous populations. The Australian COPD-X tool provides stepwise recommendations for the use of airway directed inhaled pharmacotherapy, such as short-acting BD, short-acting muscarinic antagonists, long-acting β-agonists, long-acting muscarinic antagonists, and inhaled corticosteroids (ICS) as per the severity assessed on spirometry criteria. In the current study, most patients (83%) would be classified to have either moderate or severe COPD, and therefore, are likely to be considered prescribed with ICS containing inhaled pharmacotherapy. Among the Indigenous Australian population, however, particularly in the NT, comorbid bronchiectasis and COPD is highly prevalent, and results in typically worse spirometric patterns compared with Indigenous patients with COPD in isolation. In the current study out of the 253 patients with COPD identified by spirometry, 56 (22%) had radiological evidence of concomitant bronchiectasis. Furthermore, a recent study from our centre has shown significantly higher rates of excessive yearly FEV₁ decline among Indigenous patients on ICS, which may be due to higher prevalence of comorbid bronchiectasis. Thus, the use of the COPD-X tool in this setting with no validated reference values along with higher prevalence of rather complex and concomitant respiratory comorbidities has the potential to result in overprescription of ICS, which could be detrimental to patients outcomes.

Worldwide the burden of COPD is noted to be increasingly prevalent across various ethnic and socioeconomic groups. This is also indeed true for Indigenous populations, especially among those living in high-income countries, such as the First Nation Indigenous people of the USA and Canada, Indigenous Australians and New Zealand Māori. Despite evidence in the literature suggesting that adult Indigenous people have lower spirometry values compared with their non-Indigenous counterparts, including a report by de Hamel and Glass dating back to 1975 comparing LFPs among New Zealand Māori and people of European descent, there appears to be an unprecedented gap in developing guidelines and recommendations in classifying AFL in the overall management COPD among Indigenous people globally. Although there is continued debate regarding which reference norms to use in specific populations, in the Australian context it is recommended to use GLI-2012 ‘others/mixed’ reference norms for the adult Indigenous population.

This study has demonstrated the potential effects of using these recommendations for the classification of...
AFL severity among NT Australian Indigenous patients with COPD. We observed that 75% of patients would be classified to have ‘severe’ or ‘very severe’ AFL by adopting these recommended criterion. It was outside the scope of this study to assess AFL categorisation alongside patients’ subjective experience of the disease, however, given the study sample is drawn from a referred population one may expect skewing towards the moderate/severe categories.

Although the GLI-2012 categorisation does not recommend specific treatment/management options based on AFL category, the primary care provider is likely to be heavily influenced by this and may consider different treatment options if a patient is noted to have ‘severe’ or ‘very severe’ AFL compared with ‘mild/moderate’. Underdiagnosis or overdiagnosis of COPD is not uncommon in clinical practice even in non-Indigenous patients, including misclassification and inappropriate use of airway directed inhaled pharmacotherapy.48–51 In this study, we observed that the agreement between clinical diagnosis of COPD as per medical records entry

Table 4 Clinical characteristic according to different airflow obstruction classifications

| COPD-X* | Mild (n=35) | Moderate (n=96) | Severe (n=102) | Very severe |
|---------|------------|----------------|----------------|-------------|
| Sex     | Male       | 18 (51%)       | 49 (51%)       | 57 (56%)    | –           |
| Age     | Years (mean (95% CI)) | 57.11 (53.01 to 61.21) | 56.12 (53.51 to 58.74) | 52.59 (60.53 to 54.65) | –           |
| BMI     | kg/m² (median (IQR)) | 24.54 (19.53–28.35) | 24.64 (20.94–29.66) | 21.19 (17.06–24.84) | –           |
| Smoking status | Never smoker | 1 (3%) | 12 (13%) | 5 (5%) | – |
| | Former smoker | 10 (29%) | 41 (43%) | 43 (42%) | – |
| | Current smoker | 24 (69%) | 43 (45%) | 54 (53%) | – |
| Smoking pack years (median (IQR)) | 30 (15.5–43.75) | 12.5 (5.38–40) | 26.35 (11.13–39.38) | – |
| COPD    | Clinical history of COPD | 25 (71%) | 59 (64%) | 76 (77%) | – |
| Mortality | Deceased (all cause) | 2 (6%) | 10 (10%) | 26 (25%) | – |
| | Deceased age (years) (mean (95% CI)) | 62.39 to 75.3 | 65.27 (57.02 to 73.52) | 54.07 (49.98 to 58.15) | – |

GOLD

| Mild (n=5) | Moderate (n=78) | Severe (n=97) | Very severe (n=58) |
|-----------|----------------|---------------|------------------|
| Sex       | Male           | 3 (60%)       | 41 (53%)         | 52 (54%) |
| Age       | Years (mean (95% CI)) | 52.02 (27.88 to 76.17) | 55.94 (53.14 to 58.74) | 55.88 (53.43 to 58.32) | 51.16 (48.53 to 53.78) |
| BMI       | kg/m² (median (IQR)) | 27.6 (22.27–38.57) | 24.5 (20.28–28.35) | 23.28 (19.08–28.97) | 21.74 (17.01–24.54) |
| Smoking status | Never smoker | 2 (40%) | 8 (10%) | 7 (7%) | 3 (5%) |
| | Former smoker | 1 (20%) | 22 (28%) | 47 (48%) | 25 (43%) |
| | Current smoker | 2 (40%) | 48 (62%) | 43 (44%) | 30 (52%) |
| Smoking pack years (median (IQR)) | 11.25 (4.5–112.5) | 23 (8–45) | 18 (5.1–37.5) | 27.5 (14.7–38.75) |
| COPD    | Clinical history of COPD | 1 (20%) | 50 (64%) | 66 (70%) | 44 (76%) |
| Mortality | Deceased (all cause) | 0 (0%) | 6 (8%) | 15 (15%) | 17 (29%) |
| | Deceased age (years) (mean (95% CI)) | 66.05 (52.67 to 79.42) | 62.54 (57.19 to 67.9) | 50.69 (46.1 to 55.28) |

GLI-2012

| Mild (n=16) | Moderate (n=43) | Severe (n=83) | Very severe (n=96) |
|-----------|----------------|---------------|------------------|
| Sex       | Male           | 9 (56%)       | 19 (44%)         | 44 (53%) |
| Age       | Years (mean (95% CI)) | 55.41 (47.52 to 63.29) | 57.85 (53.86 to 61.83) | 54.53 (51.98 to 57.07) | 48.35 (46.31 to 50.39) |
| BMI       | kg/m² (median (IQR)) | 23.64 (21.2–27.8) | 24.54 (20.2–28.98) | 23.99 (19.61–29.02) | 22.03 (17.21–25.81) |
| Smoking status | Never smoker | 2 (13%) | 3 (7%) | 6 (7%) | 6 (6%) |
| | Former smoker | 4 (25%) | 18 (42%) | 37 (45%) | 39 (41%) |
| | Current smoker | 10 (63%) | 22 (51%) | 40 (48%) | 51 (53%) |
| Smoking pack years (median (IQR)) | 33 (11.25–60) | 21 (8–37.5) | 12.5 (5–40) | 20 (7–37.5) |
| COPD    | Clinical history of COPD | 8 (50%) | 32 (74%) | 50 (62%) | 66 (72%) |
| Mortality | Deceased (all cause) | 1 (6%) | 3 (7%) | 10 (12%) | 21 (22%) |
| | Deceased age (years) (mean (95% CI)) | 75.3 | 62.15, 62.39 to 85.35 | 64.12 (57.43 to 70.8) | 51.14 (47.11 to 55.18) |

*Five patients excluded from COPD-X as FEV1 was >80%.

BMI, body mass index; COPD, chronic obstructive pulmonary disease; COPD-X, concise tool for chronic obstructive pulmonary disease; FEV1, forced expiratory volume in 1 s; GLI, global lung function initiative; GOLD, global initiative for chronic obstructive lung disease.
against spirometry criteria was only about 54%. It is not clear, given the fact that Indigenous patients have a higher prevalence of smoking and respiratory conditions, if health practitioners may have a preconceived perception that any or all Indigenous patients presenting with respiratory symptoms will have COPD. This further emphasises the desperate need to advocate for clinical, spirometric diagnostic and severity criteria classification, and management guidelines for COPD among the Indigenous Australian population.

The authors do acknowledge that there are only a limited number of spirometry studies available which

| Table 5  | Spirometry parameters according to different recommended classifications of airflow limitation severity |
|----------|--------------------------------------------------------------------------------------------------|
| **Mild** | COPD-X (n=35) | GOLD (n=5) | GLI-2012 (n=16) | P value |
| FVC post-BD (L) | 2.98 (2.72, 3.24) | 4.08 (2.67, 5.48) | 3.5 (2.96, 4.03) | 0.034* |
| FVC post-BD (%) | 79.51 (74.24, 84.79) | 107.8 (88.78, 126.82) | 93.19 (85.31, 101.06) | 0.003* |
| FEV₁ post-BD (%) | -1.09 (-1.31, -0.86) | 1.18 (-0.28, 2.64) | 0.07 (-0.52, 0.66) | <0.001* |
| FEV₁ post-BD (%) | 67.63 (65.77, 69.49) | 92.6 (77.53, 107.67) | 78.63 (71.78, 85.47) | <0.001* |
| FEV₁ post-BD (%) | -2.06 (-2.24, -1.88) | -0.23 (-1.18, 0.73) | -1.17 (-1.59, -0.75) | <0.001* |
| FEV₁/FVC post-BD (AV) | 0.64 (0.62, 0.66) | 0.68 (0.65, 0.7) | 0.64 (0.61, 0.68) | 0.644 |
| FEV₁/FVC post-BD (%) | 83 (80.48, 85.52) | 85.6 (78.29, 91.91) | 84.38 (80.51, 88.24) | 0.479 |
| FEV₁/FVC post-BD (%) | -2.34 (-2.57, -2.1) | -1.91 (-2.53, -1.29) | -2.3 (-2.62, -1.98) | 0.737 |

| **Moderate** | COPD-X (n=102) | GOLD (n=76) | GLI-2012 (n=43) |
| FVC post-BD (L) | 2.3 (2.16, 2.43) | 2.75 (2.58, 2.91) | 2.61 (2.39, 2.83) | 0.002 |
| FVC post-BD (%) | 62.61 (60.67, 64.56) | 72.97 (70.01, 75.94) | 73.49 (70.64, 76.34) | <0.001* |
| FVC post-BD (%) | -2.54 (-2.69, -2.38) | -1.66 (-1.84, -1.47) | -1.66 (-1.81, -1.5) | <0.001* |
| FEV₁ post-BD (L) | 1.4 (1.31, 1.48) | 1.75 (1.64, 1.85) | 1.66 (1.5, 1.82) | <0.001* |
| FEV₁ post-BD (%) | 48.46 (47.39, 49.53) | 59.83 (57.99, 61.68) | 59.12 (57.69, 61.44) | <0.001* |
| FEV₁ post-BD (%) | -3.32 (-3.42, -3.21) | -2.57 (-2.72, -2.43) | -2.54 (-2.64, -2.45) | <0.001* |
| FEV₁/FVC post-BD (AV) | 0.61 (0.59, 0.62) | 0.63 (0.62, 0.64) | 0.62 (0.6, 0.65) | 0.111 |
| FEV₁/FVC post-BD (%) | 76.91 (74.37, 79.44) | 80.23 (74.76, 83.01) | 80.47 (78.04, 82.89) | 0.054 |
| FEV₁/FVC post-BD (%) | -2.68 (-2.85, -2.5) | -2.43 (-2.59, -2.27) | -2.43 (-2.62, -1.98) | 0.042* |

| **Severe** | COPD-X (n=107) | GOLD (n=97) | GLI-2012 (n=83) |
| FVC post-BD (L) | 1.74 (1.64, 1.83) | 2.03 (1.92, 2.14) | 2.31 (2.16, 2.45) | <0.001* |
| FVC post-BD (%) | 45.95 (43.79, 48.11) | 55.8 (53.96, 57.65) | 60 (57.39, 62.61) | <0.001* |
| FVC post-BD (%) | -4.06 (-4.26, -3.86) | -3.14 (-3.31, -2.98) | -2.71 (-2.86, -2.56) | <0.001* |
| FEV₁ post-BD (L) | 0.85 (0.8, 0.9) | 1.15 (1.08, 1.21) | 1.37 (1.28, 1.47) | <0.001* |
| FEV₁ post-BD (%) | 28.27 (26.87, 29.68) | 39.99 (38.86, 41.12) | 45.1 (43.13, 47.06) | <0.001* |
| FEV₁ post-BD (%) | -4.67 (-4.8, -4.54) | -3.87 (-3.98, -3.75) | -3.49 (-3.56, -3.42) | <0.001* |
| FEV₁/FVC post-BD (AV) | 0.5 (0.48, 0.52) | 0.57 (0.55, 0.59) | 0.59 (0.58, 0.61) | <0.001* |
| FEV₁/FVC post-BD (%) | 61.85 (59.28, 64.43) | 72.38 (70.21, 74.55) | 73.72 (70.41, 77.04) | <0.001* |
| FEV₁/FVC post-BD (%) | -3.95 (-4.13, -3.78) | -3.13 (-3.32, -2.95) | -2.88 (-3.05, -2.71) | <0.001* |

| **Very severe** | COPD-X (n=58) | GOLD (n=96) |
| FVC post-BD (L) | 1.57 (1.45, 1.68) | 1.75 (1.65, 1.85) | 0.024* |
| FVC post-BD (%) | 40.97 (38.29, 43.64) | 46.79 (44.59, 48.99) | 0.001* |
| FVC post-BD (%) | -4.51 (-4.75, -4.27) | -4.19 (-4.38, -4.01) | 0.040* |
| FEV₁ post-BD (L) | 0.72 (0.66, 0.77) | 0.89 (0.82, 0.96) | <0.001* |
| FEV₁ post-BD (%) | 23.4 (22, 24.79) | 29.41 (27.73, 31.09) | <0.001* |
| FEV₁ post-BD (%) | -5.02 (-5.16, -4.88) | -4.8 (-4.91, -4.69) | 0.015* |
| FEV₁/FVC post-BD (AV) | 0.47 (0.44, 0.49) | 0.51 (0.49, 0.53) | 0.012* |
| FEV₁/FVC post-BD (%) | 57.21 (53.66, 60.75) | 63.11 (60.56, 65.67) | 0.007* |
| FEV₁/FVC post-BD (%) | -4.28 (-4.5, -4.06) | -3.94 (-4.14, -3.74) | 0.027* |

*Statistically significant.
AV, absolute value; BD, bronchodilator; COPD, chronic obstructive pulmonary disease; COPD-X, concise tool for chronic obstructive pulmonary disease; FEV₁, forced expiratory volume in 1 s; FVC, forced vital capacity; GLI, global lung initiative; GOLD, global initiative for chronic obstructive lung disease; Z, z-score.
have included Indigenous population, and that these studies may be subject to biases in participant recruitment.\textsuperscript{12-14 18 19 22 30 31 35 52–59} It is also important to acknowledge that smoking rates and pre-existing lung conditions are also highly prevalent in this population, hence recruiting study participants without smoking history or pre-existing lung disease imposes substantial challenges to conduct adult spirometry studies in an Indigenous population. Moreover, a significant proportion of Indigenous people live in regional and remote settings. Previous recommendations have suggested the use of approximate conversion factors when utilising European reference sets for other ethnic groups (ie, Hong Kong Chinese 1.0; Polynesian 0.9; North Indian and Pakistani 0.9; Japanese American 0.89; South Indian and African descent 0.87).\textsuperscript{60}

As mentioned above, the limited data available in the literature suggest that the Australian Indigenous population could have up to 20% to 30% lower values for FVC and FEV\textsubscript{1} values compared with the Australian Caucasians.\textsuperscript{12 13} Hence, it may be very reasonable to be inquisitive to know if such correctional factors could also be appropriate in this Indigenous population. Although we did not include this aspect in our main study outcome, we did create a subanalysis applying an arbitrary inflation of 15% to FVC and FEV\textsubscript{1} values (on the basis of previous research in this population demonstrating FVC and FEV\textsubscript{1} values consistently 15%-20% lower than Caucasian patients\textsuperscript{12 13 18 19 }in order to understand how the AFL severity classification would be effected with this correction factor applied (online supplemental appendix 2). In this model, instead of 238 patients being identified with COPD by FEV\textsubscript{1}/FVC <0.7 and FEV\textsubscript{1}/FVC <LLN, respectively, only 197 patients fit these criteria. Furthermore, we observed that by COPD-X and GOLD categorisation, most individuals (81% and 67%, respectively) will be re-categorised to either mild or moderate AFL; a much closer approximation of AFL severity distributions observed in non-Indigenous populations. Under ‘mild’ and ‘moderate’ AFL categorisation post-BD, LFPs significant differed between patients categorised according to COPD-X, GOLD or GLI-2012. LFPs according to COPD-X AFL categorisation were consistently one severity step removed from GOLD and GLI-2012—that is, ‘mild’ under COPD-X better matched ‘moderate’ under GOLD or GLI-2012, and ‘moderate’ under COPD-X better matched ‘severe’ under GOLD or GLI-2012.

Stakeholders, relevant organisations, such as Australian and New Zealand Society of Respiratory Science and policy-makers should consider taking into consideration of the available evidence, including realistic limitations and challenges and attempt to incorporate AFL/COPD severity classification, including considering correctional factors, to aid in the overall management of COPD in Indigenous populations.

\[\begin{align*}
\text{Figure 1} & \quad \text{Means (95% CIs) of percentage predicted values and Z-scores for postbronchodilator values by airflow limitation severity category with p values obtained via ANOVA. ANOVA, analysis of variance; COPD-X, concise tool for chronic obstructive pulmonary disease; GOLD, global initiative for chronic obstructive lung disease; FEV}_1, \text{ forced expiratory volume in 1 s; FVC, forced vital capacity; GLI, global lung function initiative.}
\end{align*}\]
this underprivileged Indigenous population. Moreover, in the authors view, health professionals caring for Indigenous patients with COPD should be made aware of potential consequences when adopting existing spirometry AFL classifications, established in non-Indigenous populations. Indeed, the context of pulmonary disease among Indigenous Australians is more similar to that of Asian populations than Caucasian populations. Including a high prevalence of concomitant chronic lung disease alongside COPD, in particular the presence of bronchiectasis. The position statement of the Asian Pacific Society of Respirology suggests cautious use of ICS in their population.

Our study, for the first time, has highlighted the impact of adopting particular AFL criterion for the Australian NT Indigenous population with COPD. Hence, this study is of significant clinical relevance in addressing and highlighting this importance of COPD severity classification and the use of spirometry reference ranges among Indigenous patients with COPD. Health professionals, especially primary health practitioners in regional and rural/remote communities caring for Indigenous patients, should be aware of potential implications in classifying AFL severity for clinical decision making, in particular, in the Australian context while using COPD-X recommendations. This study also highlights the importance for healthcare providers for Indigenous people not swapping between various COPD severity classifications which would lead to inconsistencies. Moreover, it is also vital to understand that reference values and guidelines developed for other ethnic populations, including Caucasians cannot be considered as ‘norm’ for Indigenous people and the approach to the holistic management of COPD should be different to their Caucasian counterpart. As such, spirometry plays an integral role in the diagnosis, quantifying the severity, and management of COPD with inhaled pharmacotherapy according to the severity classification. Although significant progress has been made towards prevention and management of COPD in Australia and globally, within the Indigenous Australian population the problem still remains - more specifically, in relation to establishing COPD diagnosis/severity classification and management guidelines. Further dedicated efforts are required to close the respiratory health gap in our Indigenous population, not only in reducing COPD related morbidity and mortality, but also in reducing healthcare cost and utilisation.

**Limitations**

The results of this study are restricted to the Australian Indigenous study participants from the TEHS region of the NT of Australia and the outcomes represented in this study may or may not be applicable to other Indigenous populations globally. Moreover, in the lack of similar studies in other non-Australian Indigenous cohort we were unable to compare if our study outcomes compare to other Indigenous populations. Nevertheless, this is the first ever study to provide insight into the implications of utilising the established AFL classifications in an Indigenous population, and we anticipate this study will be of interest and be influential among researchers globally to address these important issues for their respective Indigenous populations. The authors are not aware of any updated AFL/COPD recommendations or guidelines in any Indigenous population globally at the time of writing of this paper. Other potential limitations that may or may not be relevant specifically to this current study are described in our previously published reports.

**CONCLUSION**

Irrespective of which of the three recommended AFL classifications used (COPD-X, GOLD, GLI-2012), most Indigenous patients with COPD referred for lung function testing will likely fall into either the severe or very severe category. Spirometry values for FVC and FEV₁ vary significantly between the three recommended criteria. In the absence of established spirometry reference norms among the adult Indigenous population, health professionals should be aware of implications in using various recommended AFL classifications in the overall management of Indigenous patients presenting with COPD, including, prescribing inhaled pharmacotherapy.

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**Contributors**

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**Competing interests**

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**Patient consent for publication**

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