Insights into Chronic Obstructive Pulmonary Disease
Epidemiology, Phenotypes and Outcomes
from SPIROMICS

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ABSTRACT

SubPopulations and InteRmediate Outcome Measures in COPD Study (SPIROMICS) is a multicentre National Institutes of Health funded cohort study of nearly 3000 never-smokers and ever-smokers (≥ 20 pack-years) with and without chronic obstructive pulmonary disease (COPD). It was designed to help fill gaps in our understanding of the biological complexity and clinical heterogeneity of COPD. The original goals for this study were to identify subgroups of smokers who may benefit from targeted therapies and to discover and validate intermediate biomarker endpoints. Here we review important findings from SPIROMICS over the past five years, including the impact of environmental and occupational exposures on the respiratory health of smokers, the characterisation of symptomatic smokers with preserved pulmonary function and the investigation of several diagnostic and prognostic biomarkers such as airway mucin concentration, functional small airways disease on chest computed tomography and a gene expression signature of interleukin 17A response in airway epithelium. (BRN Rev. 2019;5(4):233-48)

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INTRODUCTION

Chronic obstructive pulmonary disease (COPD) affects millions of people worldwide and is associated with high morbidity and mortality. The complexity of COPD extends far beyond incompletely reversible airflow limitation as reflected by the striking heterogeneity in disease presentation, progression and outcomes. Therefore, the characterisation of distinct clinical phenotypes is paramount to guide the promising field of precision medicine in COPD. SubPopulations and InteRmediate Outcome Measures in COPD Study (SPIROMICS) is a multicentre, National Institutes of Health (NIH)-funded cohort study of nearly 3000 never-smokers and ever-smokers (≥ 20 pack-years) with and without COPD. Table 1 shows the demographic, anthropometric and clinical characteristics of SPIROMICS participants at baseline. The SPIROMICS study was designed to help fill gaps in our understanding of the clinical heterogeneity and biological complexity of COPD. The original primary aims were to 1) identify subgroups of smokers who may eventually benefit from specific therapies as part of targeted clinical trials, and 2) discover and validate biomarkers to serve as intermediate outcomes in such trials. In this review article, we outline important studies from SPIROMICS that significantly advanced our knowledge of COPD epidemiology, phenotypes and outcomes.

EXPOSURES

While active smoking is a strong risk factor for COPD development and a well-established predictor of disease progression, the impact of passive smoke exposure, so-called “second-hand smoke”, on COPD outcomes has been unclear. In an analysis of 1580 participants with COPD, 27% reported living with a smoker and 20% were exposed to second-hand smoke during the past week. Living with a smoker was associated with worse respiratory-related quality of life as measured by the St. George’s Respiratory Questionnaire (SGRQ), a higher burden of respiratory symptoms as measured by the COPD Assessment Test (CAT) and a higher incidence of severe COPD exacerbations. Similarly, exposure to second-hand smoke in the past week was associated with worse respiratory symptoms, including chronic productive cough (odds ratio [OR] 1.77; 95% confidence intervals [CI] 1.33-2.35) and wheezing (OR 1.34; 95% CI 1.02-1.77), even after adjusting for demographics, lung function, personal smoking status and smoking history. Interestingly, current smokers, obese individuals and participants with Global Initiative for Chronic Obstructive Lung Disease (GOLD) 1-2 grades of spirometry severity were most affected. Therefore, this study not only characterises the harms of passive smoke exposure among individuals with COPD but also identifies subgroups most susceptible to its effects.

Electronic cigarette (e-cigarette) and marijuana use has been increasing among cigarette smokers with or at risk for COPD. The long-term impact of the consumption of such products is still being investigated. A study of 3536 Genetic Epidemiology of COPD (COPD-Gene) and 1060 SPIROMICS current or former cigarette smokers found that 12-16% of participants ever used e-cigarettes and that 5% were current users. Although close to 90% of ever-users reported using e-cigarettes in order to help decrease or quit consumption of...
regular cigarettes, these goals were not consistently reached at the 5-year follow-up visit. After adjusting for relevant clinical confounders, ever-use of e-cigarettes was associated with higher SGRQ scores, a higher prevalence of chronic bronchitis and a higher incidence of COPD exacerbations. In contrast, neither current nor former use of marijuana was associated with worse respiratory symptoms though the prevalence of marijuana ever-use was 51%\(^{10}\). Compared to never-users, current and former marijuana users were younger, had better lung function, and were more likely to be men and current smokers of cigarettes. Longitudinal studies are needed to better understand the long-term effects of e-cigarettes and marijuana on the respiratory health, lung function and clinical outcomes of individuals with COPD.

Occupational exposure to vapours, dust, gases or fumes has been recognised as an important risk factor for COPD\(^ {11}\). Paulin et al.\(^ {12}\) further confirmed this relationship as they found that SPIROMICS participants with an intermediate-to-high level of occupational exposures had 44% higher odds (95% CI 1.04-1.97) of COPD, even after adjusting for demographics and smoking history. Among individuals with COPD, the presence of such exposures was associated with a higher burden of respiratory symptoms, decreased exertional capacity and a higher incidence of moderate and severe COPD exacerbations. Furthermore, occupational exposures were also associated with radiographic abnormalities on chest computed tomography (CT) including airways disease, air trapping and emphysema\(^ {13}\). These findings highlight the importance of obtaining a thorough occupational history both in individuals with newly diagnosed and previously established COPD. Outcomes may also be affected by place of residence. In fact, rural residence has been independently associated with an increased risk of moderate COPD exacerbations\(^ {14}\). Accounting
for agricultural occupation attenuated but did not fully explain this association, highlighting the need for further investigations into the triggers and mechanisms of COPD exacerbations in residents of rural areas.

**SMOKERS WITHOUT AIRFLOW LIMITATION**

While a post-bronchodilator forced expiratory volume in the first second (FEV$_1$)/forced vital capacity (FVC) ratio < 0.70 is currently needed to make a diagnosis of COPD$^{15}$, many smokers who do not meet this spirometric definition still experience substantial morbidity. A landmark SPIROMICS study showed that half of ever-smokers with preserved pulmonary function reported a significant burden of respiratory symptoms as determined by a CAT score $\geq$ 10$^{16}$. These symptomatic smokers with preserved pulmonary function had a higher rate of respiratory exacerbations than their asymptomatic counterparts (0.27 ± 0.67 versus 0.08 ± 0.31 events per year, $p < 0.001$) (Fig. 1). This association persisted in adjusted analyses and when cut-offs of 0.65, 0.70 and the lower limit of normal (LLN) were used for FEV$_1$/FVC to define obstruction. Notably, close to half of these symptomatic current or former smokers were using bronchodilators and close to a quarter of them were on inhaled corticosteroids, even though no evidence basis for the use of such medications in this patient population exists yet. This analysis helped fulfil one of the primary aims of SPIROMICS, namely to identify a subgroup of patients for targeted enrolment in clinical trials, as it provided the scientific basis for the currently enrolling REdefining THERapy IN Early COPD (RETHINC) trial for the Pulmonary Trials Cooperative (clinicaltrials.gov identifier: NCT02867761). RETHINC is a multicentre, randomised, placebo-controlled trial testing the efficacy of the combined long-acting beta agonist/long-acting muscarinic antagonist (LABA/LAMA) inhaler indacaterol/glycopyrrrolate in symptomatic smokers without airflow obstruction. The primary outcome is the proportion of individuals with a 4-point improvement in SGRQ score at 12 weeks.

The cut-off used to identify the presence of an obstructive ventilatory defect on spirometry (FEV$_1$/FVC < 0.70 versus < LLN) remains a controversial topic. Pirozzi et al.$^{17}$ sought to understand the characteristics of ever-smokers with FEV$_1$/FVC < 0.70 but > LLN. Compared to ever-smokers and never-smokers with FEV$_1$/FVC > 0.70, this discordant group had a lower post-bronchodilator FEV$_1$ and a greater extent of both emphysema and non-emphysematous air trapping on chest CT. In fact, 44% of participants in this group had at least one of these radiographic abnormalities. Compared with an FEV$_1$/FVC cut-off set at the LLN, a cut-off of 0.70 had a higher sensitivity (85% versus 78%) but a lower specificity (72% versus 81%) for identifying smokers with radiographic emphysema or gas trapping. This is yet another study that highlights the drawbacks of relying on lung function alone to diagnose COPD as many smokers with normal spirometry have a significant burden of clinical and radiographic disease.

**CHEST IMAGING**

Chest CT has become a widely available imaging modality that is increasingly ordered...
in the inpatient and outpatient settings to answer a number of clinical questions ranging from ruling out a pulmonary embolism to screening for lung cancer. Beyond providing the requested clinical information, chest CTs contain a wealth of additional data on pulmonary structures (airways, lung parenchyma, pulmonary vasculature), and can also inform comorbidities such as coronary artery disease through coronary artery calcium scoring, osteoporosis through vertebral bone mineral density and frailty through pectoralis muscle area measurement\(^{18,19}\). Chest CT is an integral component of the clinical phenotyping of SPIROMICS participants with rigorous imaging protocols implemented through the quantitative CT lung assessment system (QCT-LAS)\(^{20}\).

To limit the bias of airway selection, Smith et al.\(^{21}\) compared spatially matched airways on chest CTs of participants with COPD and controls enrolled in SPIROMICS and the Multi-Ethnic Study of Atherosclerosis (MESA). They found airway wall areas for central airway segments (generations 1-6 in SPIROMICS and 3-6 in MESA) to be significantly smaller in COPD compared to controls and to become progressively smaller with increasing GOLD grades of spirometry severity. This study underlines the importance of spatial matching...
as selecting airways at random or based on lumen diameter will result in a biased comparison of more proximal airways in COPD to more distal airways in non-COPD. In another CT analysis of the SPIROMICS and MESA cohorts, airway branch variants were found in more than a quarter of the general population. The most common airway branch variant, the presence of an accessory sub-superior airway, was found in 16.0% of participants and was associated with COPD (pooled OR 1.31, 95% CI 1.10-1.55 in adjusted analyses) and with shorter central airways and increased emphysema in all lobes. The second most common airway branch variant, the absence of a right medial-basal airway, was found in 6.1% of participants and was associated with COPD among smokers (pooled OR 1.57; 95% CI 1.14-2.17 in adjusted analyses), small airway lumen areas in all lobes and with two single nucleotide polymorphisms in the fibroblast growth factor (FGF) 10 gene. These results show that central airway branch variations may represent a heritable risk factor for COPD development and are linked to diffuse pathology in airway architecture and lung parenchyma.

Emphysema can be either visually assessed or quantitatively measured on CT. In an analysis of the SPIROMICS and COPDGene cohorts, higher extent of emphysema on CT (measured as the percentage of lung voxels < -950 Hounsfield Units [HU]) was associated with a higher incidence of exacerbations and all-cause mortality (Fig. 2). In SPIROMICS, compared with participants with < 5% emphysema, those with ≥ 5% emphysema had a higher mean exacerbation frequency (0.46 versus 0.21 exacerbations/year; p < 0.001) and higher mortality (2.66 versus 0.89 deaths per 100 person-years; p = 0.01). Results were similar in COPDGene. Therefore, extent of emphysema on an available CT can be a powerful tool to predict patient outcomes.

Coronary artery disease is a common comorbidity in COPD. In an analysis of 300 SPIROMICS participants, Bhatt et al. demonstrated that while FEV₁/FVC was associated with coronary artery calcium (assessed through the Weston visual score), quantitative emphysema and airway wall thickening on CT were not. In subsequent analyses, they found centrilobular, but not paraseptal, emphysema to be associated with coronary artery calcium. Furthermore, they demonstrated that certain plasma biomarkers of inflammation such as C-X-C motif chemokine ligand 9 (CXCL9) mediated the association between airflow obstruction and coronary artery calcium. The importance of this study is twofold: 1) it further characterises the link between COPD and coronary artery disease, and 2) it highlights the utility of visually assessing emphysema subtypes as a complementary tool to quantitatively measuring emphysema.

Parametric response mapping (PRM) is an imaging application that incorporates voxel-based changes in lung parenchyma density between co-registered inspiratory and expiratory chest CTs to classify lung as normal, emphysema (< -950 HU on inspiration and < -856 HU on expiration) and non-emphysematous air trapping (> -950 HU on inspiration and < -856 HU on expiration), a measure of functional small airways disease (fSAD). Haghighi et al. identified four different chest imaging clusters among 284 current smokers from SPIROMICS. Cluster 1 was characterised by normal airways and minimal fSAD and emphysema, cluster 2 by airway wall thickening and increased fSAD (mean 8.4%), cluster 3 by even higher...
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fSAD (mean 12.3%) but no airway wall thickening, and cluster 4 by the highest fSAD (mean 34.9%) and emphysema (13.5%). As opposed to cluster 1, which had preserved pulmonary function, the other clusters were characterised by airflow obstruction of increasing severity (mean FEV₁ % predicted of 0.80, 0.76 and 0.49 in clusters 2, 3 and 4, respectively). Cluster 4 had the highest mean CAT score and the highest rate of exacerbations. Although this is a smaller study including current smokers only, it highlights the relationship between various CT metrics and clinical variables. Expanding on this relationship, Martinez et al.²⁹ found a positive association between fSAD and each of age and FVC, but not FEV₁, in a cross-sectional analysis of participants without airflow obstruction. These results shed light on the complex link between ageing and respiratory physiologic changes, and also underscore the potential of PRM for phenotyping individuals without airflow limitation on spirometry. Beyond cross-sectional analyses, PRM metrics can serve as longitudinal imaging biomarkers. Mathematical model simulations showed that fSAD may be a transitional pathologic phase between normal lung and emphysema³⁰. Subsequent 5-year longitudinal analyses from COPDGene supported this hypothesis³¹. In summary, these remarkable advances in chest imaging have improved our understanding of COPD and have helped characterise some of its clinical phenotypes. It remains to be seen how they can best be incorporated within routine clinical care and whether they can be leveraged to detect early disease.

Figure 2. (A) Plot of mean exacerbations per year by percentage emphysema in SPIROMICS. Black dots represent raw averages for exacerbation rates grouped by nearest emphysema percentile. Age-adjusted estimates for average exacerbations per year are shown via linear splines in black and smoothed splines in red. (B) Plot of average deaths per 100 person-years by percentage emphysema in SPIROMICS. Black dots represent raw averages for death rates grouped by nearest emphysema percentile. Age-adjusted estimates for average deaths per 100 person-years are shown via linear splines in black and smoothed splines in red (reproduced from Han MK²⁵ with permission of the American Thoracic Society, © 2019 American Thoracic Society).
BIOMARKERS

Complete blood counts are frequently obtained in clinical practice and are readily available for many patients. Both normocytic anaemia and thrombocytosis were found to be independently associated with worse dyspnoea and respiratory-related quality of life in individuals with COPD32,33. In addition, thrombocytosis was linked to a higher likelihood of all COPD exacerbations (adjusted OR 1.5; 95% CI 1.1-2.0) and severe COPD exacerbations (adjusted OR 1.5; 95% CI 1.1-2.2) during the preceding year in a combined analysis of the SPIROMICS and COPDGene cohorts33. Interestingly, in an observational propensity-score matched analysis from SPIROMICS, aspirin users experienced fewer exacerbations (incidence rate ratio [IRR] 0.78, 95% CI 0.65-0.94) and had a lower burden of respiratory symptoms34. These findings deserve further validation in a randomised controlled trial.

The clinical utility of the blood and sputum eosinophil counts in patients with COPD remains controversial35. SPIROMICS participants with elevated sputum eosinophils (≥ 1.25%) had worse lung function, more emphysema and small airways disease on CT and a higher rate of severe exacerbations during the preceding year compared with participants with lower sputum eosinophils (< 1.25%)36. While participants with high blood eosinophils (≥ 200 cells/µL) had worse lung function and higher airway wall thickness than those with lower blood eosinophils (< 200 cells/µL), there were no significant differences between these groups with regards to history of exacerbations and evidence of radiographic emphysema and small airways disease. Although the association between blood and sputum eosinophil counts was statistically significant, it remained relatively weak (area under the curve [AUC] 0.64). Therefore, sputum eosinophils appears to be a better biomarker of disease severity and morbidity than blood eosinophils. Broncho-alveolar lavage (BAL) is another compartment where eosinophil count can be measured. Among individuals with COPD, current smokers had significantly higher BAL eosinophils than former smokers37. In contrast, current and former smokers without COPD had similar BAL eosinophil counts. These results suggest that smoking status affects eosinophil recruitment and/or retention in the COPD lung. Of note, BAL eosinophil count was correlated with neither blood nor sputum eosinophil counts.

The generalised and local inflammatory milieu in COPD is reflected in part by disturbed levels of multiple cytokines and chemokines. For example, plasma interleukin 6 (IL-6), a pro-inflammatory cytokine, has been associated with airflow obstruction as well as lung function decline and progression of emphysema over five years of follow-up38. Another cytokine of interest is IL-17A whose effects result in the recruitment and activation of neutrophils and macrophages in the airway epithelium39. After generating and validating a gene expression signature of IL-17A response in bronchial airway epithelial brushings, Christenson et al.40 found this signature to be associated with lower lung function and higher radiographic fSAD in ever-smokers with COPD. When this IL-17A gene expression signature was tested in the Groningen and Leiden Universities study of Corticosteroids in Obstructive Lung Disease (GLUCOLD) study which randomised participants to inhaled corticosteroids (ICS) versus placebo, it was associated
with decreased response to ICS independent of airway eosinophilic or type 2 inflammation. Therefore, this biomarker identifies a subgroup of COPD patients who may not benefit from traditional ICS therapy but may rather respond to better targeted anti-inflammatory agents. Measurement of serum inflammatory biomarkers may also have a role in smokers without COPD as higher levels of C-reactive protein and soluble tumour necrosis factor (TNF) receptors were associated with a higher burden of respiratory symptoms and a lower 6-minute walking distance in this patient population.

Many patients with COPD experience chronic bronchitis defined as cough with sputum expectoration for at least three months a year during two consecutive years. Airway mucin concentration as measured by size-exclusion chromatography and refractometry appears to be a valid biomarker of chronic bronchitis. It was significantly increased in participants who met the clinical definition of chronic bronchitis compared to those who did not (Fig. 3), in participants with severe COPD compared to never-smoker controls, and in participants with ≥2 respiratory exacerbations per year compared to those with none. Furthermore, higher airway mucin concentration was associated with lower forced expiratory flows in the mid-expiratory phase (FEF25-75%) on spirometry and with higher fSAD and relative airway wall thickness on CT.

The quest to find reliable biomarkers predictive of respiratory exacerbations has proven challenging. In an analysis of participants from SPIROMICS and COPDGene assessing the predictive ability of 90 different serum or plasma proteins, there was minimal replication between the two cohorts. This may be related to the stochastic and highly variable nature of exacerbation events. In an analysis of 1843 SPIROMICS participants with COPD, 49% experienced at least one exacerbation during three years of follow-up with only 7% experiencing at least one exacerbation during each year and 2% experiencing ≥2 exacerbations during each year (Fig. 4). Instead, an inconsistent pattern of exacerbation occurrences characterised by years with and years without exacerbations was frequent. Another factor complicating the identification of reliable biomarkers to

![Figure 3. Box plots of total airway mucin concentration in current or former smokers who were identified as having chronic bronchitis by a questionnaire of its classic definition (199 participants), current or former smokers who were identified as having chronic bronchitis by the St. George’s Respiratory Questionnaire (SGRQ) (382 participants), and controls who had never smoked and were not identified as having chronic bronchitis by either questionnaire. P values are for the comparison with healthy controls who had never smoked (reproduced from Kesimer M, © 2017 Massachusetts Medical Society; reprinted with permission from Massachusetts Medical Society).](image-url)
predict COPD exacerbations is the effect of common genetic polymorphisms on the measurement of blood proteins as demonstrated in a combined analysis of SPIROMICS and COPDGene\textsuperscript{46}.

Despite these challenges, a few serum biomarkers were found to be independently associated with incident COPD exacerbations, including low immunoglobulin A (IgA), high N-terminal pro-brain natriuretic peptide (NT-proBNP), and high matrix metalloproteinase 9 (MMP-9). Putcha et al.\textsuperscript{47} found a serum IgA level \(\leq 70\) mg/dL to be associated with a higher incidence of exacerbations (IRR 1.71; 95\% CI 1.01-2.87; \(p = 0.04\)) after adjusting for relevant clinical covariates\textsuperscript{47}. In a separate SPIROMICS analysis, participants in GOLD grade D (high symptom and exacerbation burdens) had higher serum NT-proBNP levels compared with participants in GOLD grade B (high symptom but low exacerbation burdens) (758.4 versus 574.9 pg/ml, \(p = 0.03\))\textsuperscript{48}. Further, an NT-proBNP level \(\geq 900\) pg/ml was independently associated with an increased incidence of COPD exacerbations (IRR 1.62; 95\% CI 1.19-2.21; \(p = 0.002\)). This association was maintained in participants with and without a self-reported history of coronary artery disease, myocardial infarction and congestive heart failure. In adjusted models, Wells et al.\textsuperscript{49} found an association between elevated plasma MMP-9 (defined as > 95\% percentile of values from healthy controls) and increased odds of exacerbations (OR 1.71; 95\% CI 1.00-2.90; \(p = 0.049\)), frequency of exacerbations (IRR 1.45; 95\% CI 1.23-1.70; \(p < 0.001\)) and shorter time to first exacerbation (21.7 versus 31.7 months; \(p = 0.015\)). Importantly, these results were replicated in participants

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\caption{Frequency of acute exacerbations in each of the three years in patients with chronic obstructive pulmonary disease. Data are the proportion of patients with each category of acute exacerbation frequency, by GOLD grade and in the entire group (n = 1105) (reprinted from Han MK\textsuperscript{45}, © 2017 The Lancet, with permission from Elsevier).
GOLD: Global Initiative for Chronic Obstructive Lung Disease.}
\end{figure}
from the COPDGene cohort. In summary, low IgA, high NT-proBNP and high MMP-9 likely reflect increased susceptibility to exacerbation events through impaired immunity, clinical or subclinical cardiac dysfunction and increased inflammation, respectively. It remains to be determined whether targeted therapies (such as intravenous immunoglobulin replacement or MMP-9 modulators) and interventions (such as more aggressive screening for cardiovascular disease) can reduce exacerbation frequency in select patients with COPD.

In a genome-wide association study (GWAS) of 1645 non-Hispanic white participants enrolled in SPIROMICS, a functional rare variant in the SERPINA1 gene (rs28929474: Glu342Lys) was associated with lower FEV1/FVC and FEV150. In a model including age, sex and smoking pack-years, the top 10 single nucleotide polymorphisms associated with FEV1 (including the aforementioned SERPINA1 variant) explained 8.6% of the variance of FEV1/FVC. This study provides additional evidence of the genetic contributions to COPD pathogenesis and underlines the potential of combining genetic and clinical data.

COMORBIDITIES AND QUALITY OF LIFE

Comorbidities are common in COPD and include coronary artery disease, osteoporosis, depression, anxiety and obstructive sleep apnoea. A simple count of comorbidities performed as well as weighted comorbidity scores to predict the modified medical research council (mMRC) score of dyspnoea, the SGRQ score, the 6-minute walking distance and the risk of exacerbations, even after accounting for demographics, smoking history and lung function51. Zeidler et al.52 found that poor sleep quality as measured by the Pittsburgh Sleep Quality Index was associated with worse quality of life among patients with COPD beyond their risk for obstructive sleep apnoea52. As more than half of participants reported poor sleep, intervening on this reversible problem in clinical practice has the potential to significantly improve lives of patients with COPD. Dyspnoea severity also expectedly contributes to the quality of life of individuals with COPD. However, for the same degree of dyspnoea, middle-age patients with COPD (50-64 years) have higher SGRQ scores (i.e., worse health quality of life) than their older counterparts (65-80 years), which further highlights the importance of taking into account age-related differences when delivering individualized care in everyday practice53.

INHALER THERAPY

The GOLD provides guidance on management of inhaler therapy for COPD patients based on its ABCD staging schema. Ghosh et al.54 assessed the alignment of inhaler therapy prescribed to SPIROMICS participants by their medical providers between 2010 and 2016 with 2011 GOLD recommendations54. They found that close to half of participants were not being managed in line with GOLD strategies (Fig. 5A). Of these, 54% were under-treated and 46% were over-treated (Fig. 5B). Under-treatment was mainly due to lack of long-acting inhalers in GOLD grade D, while over-treatment was mostly attributed
to the use of ICS-containing inhalers when not indicated. Both under- and over-treatment may result in higher morbidity in patients with COPD as the former may hinder the achievement of important patient-centred outcomes such as symptom relief and reduction of exacerbation frequency, while the latter may lead to unnecessary adverse events such as pneumonia and non-tuberculous mycobacteria infection. Therefore, a better understanding of the barriers limiting the widespread adoption of GOLD recommendations in clinical practice warrants further study.

CONCLUSIONS

Over the past five years, SPIROMICS has generated significant knowledge on the clinical phenotypes of COPD through the identification of distinct subgroups and the investigation of various chest and imaging biomarkers. Table 2 summarises the major findings of SPIROMICS to date and highlights important gaps in our understanding of COPD epidemiology, pathogenesis and progression. The benefits of lung-volume-reduction surgery in specific patients with emphysema and of roflumilast in frequently exacerbating individuals with chronic bronchitis have established the feasibility and potential of precision medicine in COPD. However, beyond these select therapies, most patients receive the same treatments, largely consisting of inhaled bronchodilators and corticosteroids. Therefore, therapies targeted towards specific underlying pathologic pathways remain sorely needed in COPD. To this end, SPIROMICS II will extend follow-up of previously enrolled participants with aims to define the natural
history of symptomatic smokers without airflow obstruction, elucidate the molecular phenotypes behind clinical and radiographic disease progression, and further characterise the triggers and host inflammatory responses associated with COPD exacerbations. Such knowledge will help expand our understanding of the complex heterogeneity of COPD to advance precision medicine and improve clinical outcomes.
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