Genetic variation in genes regulating skeletal muscle regeneration and tissue remodelling associated with weight loss in chronic obstructive pulmonary disease

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Abstract

Background Chronic obstructive pulmonary disease (COPD) is the third leading cause of death globally. COPD patients with cachexia or weight loss have increased risk of death independent of body mass index (BMI) and lung function. We tested the hypothesis genetic variation is associated with weight loss in COPD using a genome-wide association study approach.

Methods Participants with COPD (N = 4308) from three studies (COPDGene, ECLIPSE, and SPIROMICS) were analysed. Discovery analyses were performed in COPDGene with replication in SPIROMICS and ECLIPSE. In COPDGene, weight loss was defined as self-reported unintentional weight loss > 5% in the past year or low BMI (BMI < 20 kg/m²). In ECLIPSE and SPIROMICS, weight loss was calculated using available longitudinal visits. Stratified analyses were performed among African American (AA) and Non-Hispanic White (NHW) participants with COPD. Single variant and gene-based analyses were performed adjusting for confounders. Fine mapping was performed using a Bayesian approach integrating genetic association results with linkage disequilibrium and functional annotation. Significant gene networks were identified by integrating genetic regions associated with weight loss with skeletal muscle protein–protein interaction (PPI) data.

Results At the single variant level, only the rs35368512 variant, intergenic to GRXCR1 and LINCO02383, was associated with weight loss (odds ratio = 3.6, 95% confidence interval = 2.3–5.6, P = 3.2 × 10⁻⁸) among AA COPD participants in COPDGene. At the gene level in COPDGene, EFNA2 and BAIA2 were significantly associated with weight loss in AA and NHW COPD participants, respectively. The EFNA2 association replicated among AA from SPIROMICS (P = 0.0014), whereas the BAIA2 association replicated in NHW from ECLIPSE (P = 0.025). The EFNA2 gene encodes the membrane-bound protein ephrin-A2 involved in the regulation of developmental processes and adult tissue homeostasis such as skeletal muscle. The BAIA2 gene encodes the insulin-responsive protein of mass 53 kD (IRSp53), a negative

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regulator of myogenic differentiation. Integration of the gene-based findings participants with PPI data revealed networks of genes involved in pathways such as Rho and synapse signalling.

Conclusions The **EFNA2** and **BAIAP2** genes were significantly associated with weight loss in COPD participants. Collectively, the integrative network analyses indicated genetic variation associated with weight loss in COPD may influence skeletal muscle regeneration and tissue remodelling.

Keywords GWAS; Cachexia; Weight loss; COPD; Genetics; Biomarkers; Skeletal muscle regeneration; Tissue remodelling

**Introduction**

Chronic obstructive pulmonary disease (COPD) is the third leading cause of death internationally with associated mortality continuing to rise.1, 2 Although COPD is primarily diagnosed using lung function, traits not directly related to lung function such as cachexia greatly reduce quality of life and increase risk of mortality.3, 4 Cachexia is a debilitating comorbidity increasing risk of death and healthcare expenditure.5 Most often thought of with respect to cancer, it has been estimated that there are 1.4 times as many patients with COPD cachexia than cancer cachexia by population prevalence.6

Cachexia is defined as weight loss, primarily caused by loss of muscle with or without loss of fat, in individuals suffering from a chronic illness.3 The consensus definition for cachexia diagnosis includes weight loss > 5% in the last 12 months or low body mass index (BMI) (BMI < 20 kg/m²) in addition to three out of five of decreased muscle strength; fatigue; anorexia; low fat-free mass index; and any indication of increased inflammatory markers (C-reactive protein, IL6, etc.), anaemia, or low serum albumin.7 We recently demonstrated participants with COPD with cachexia and/or weight loss greater than 5% in the past year had a greater than three-fold increased mortality independent of BMI and lung function.4 Monitoring cachexia using weight loss criteria is relatively simple and predictive of mortality among individuals with COPD.

Loss of muscle mass underlying weight loss in cachexia can be influenced by dysregulation of a number of mechanisms involved in the balance between protein synthesis and degradation.7 Further, impaired ability to regenerate skeletal muscle tissue can contribute to muscle loss in COPD cachexia.7 COPD patients with advanced disease are more likely to exhibit weight loss4 in addition to skeletal muscle remodelling from a slow twitch (Type I) to a fast twitch (Type II) myofibre shift.8 Skeletal muscle remodelling occurs in response to external stimuli leading to activation of intracellular signalling pathways and muscle fibre transition.8

Although the major risk factor for COPD is smoking, COPD is a heritable disease with multiple genetic loci reproducibly associated.9–13 The prevalence of cachexia in COPD is correlated with increasing disease severity.9 Genetic variation may also contribute to the development of cachexia and weight loss in COPD. By performing genome-wide association study (GWAS) analyses, we previously identified genetic variants associated with longitudinal BMI in a small number of participants with COPD (N = 237) in the Framingham Heart Study.14 As analyses were performed in a small size, investigation in a larger sample of participants with COPD with more precise phenotyping is merited. The genetics of cachexia have been more thoroughly investigated in cancer with several genes reproducibly associated using primarily candidate gene association approaches.15, 16 Cancer cachexia genes identified in association studies are known to be involved with inflammatory response regulation, pathways directing muscle and fat metabolism and appetite regulation among others.15, 17

We hypothesized genetic variation may be associated with weight loss contributing to cachexia in COPD. To test this hypothesis, we performed GWAS testing in 4308 participants with COPD from 3 cohorts: COPD Genetic Epidemiology (COPDGene), Evaluation of COPD Longitudinally to Identify Predictive SurrogateEndpoints (ECLIPSE), and Subpopulations and Intermediate Outcome Measures in COPD Study (SPIROMICS). Replication was assessed in remaining cohorts. GWAS findings were further explored by integrating gene-based findings with publicly available transcriptomics and protein–protein interaction (PPI) databases to gain additional insight to underlying biological mechanisms which may be influencing cachexia.

**Materials and methods**

**Ethics statement**

Institutional Review Board approval for all analyses was obtained from the University of Alabama at Birmingham and performed in accordance with the ethical standards laid down in the 1964 Declaration of Helsinki and its later amendments. Manuscript complies with the Ethical guidelines for authorship manuscript that they comply with the Ethical guidelines for authorship and publishing in the Journal of Cachexia, Sarcopenia and Muscle.18
**Study participants**

The current analyses utilized COPD participants recruited as part of three studies: COPDGene,19 ECLIPSE,20 and SPIROMICS21 (Supporting Information, Figure S1). In all studies, COPD was classified using post-bronchodilator lung function testing (FEV1: forced expiratory volume in 1 s and FEV1/FVC: FEV1 expressed as a fraction of forced vital capacity) at baseline enrollment. All participants with COPD had moderate to severe disease defined by a GOLD22 stage of 2 (FEV1/FVC < 0.7 and 50% < FEV1 < 80% predicted) or higher (GOLD 3 and 4). In the current analyses, COPDGene COPD participants had at least a 10 pack-year smoking history, were aged 45 to 80 years at baseline and followed longitudinally with two visits 5 years apart. In ECLIPSE, COPD participants had at least a 10 pack-year smoking history and were aged 45 to 75 years at baseline and followed longitudinally. Visits in ECLIPSE occurred at baseline, 3 months, 6 months, and then every 6 months for 3 years. In SPIROMICS, COPD participants had at least a 20 pack-year smoking history, were aged 41 to 80 years at baseline, and were followed with annual visits for 3 years.

**Weight loss in chronic obstructive pulmonary disease**

As diagrammed in Figure S1, COPDGene weight loss was defined as either self-reported, unintentional weight loss greater than 5% in the past year, or as had low BMI (<20 kg/m²). We performed additional cleaning of the self-reported unintentional weight loss variable in COPDGene by confirming weight loss based on the weight measurements collected at Visits 1 and 2. This led to the exclusion of two participants (Figure S1). In ECLIPSE and SPIROMICS, weight loss greater than 5% was defined if present at any of the longitudinal visits.

**Genome-wide association study analyses**

Genotyping in COPDGene was performed using the Illumina Human Omni 1-Quad (Illumina, San Diego, CA), in ECLIPSE using the Illumina HumanHap 550v3 chips and in SPIROMICS using the Illumina HumanOmniExpressExome BeadChip (Illumina, Inc., San Diego, CA). Standard quality control steps were performed on DNA samples and single-nucleotide polymorphism (SNP) data as previously described.23–25 For all three studies, genotypes were imputed using the Haplotypereference Consortium reference panel.26 Only SNPs with an imputation quality score of 0.5 or greater were included in the analysis. SNP positions were reported on the human genome 19 build. SNPs were annotated to genes or closest genes using the NCBI human genome 19 RefGene database (version 2017-06-01) as implemented using ANNOVAR.27 In COPDGene, a total of 5,405,435 and 7,629,332 variants were imputed and passed quality control in the Non-Hispanic White (NHW) and African American (AA) COPD participants, respectively. In ECLIPSE, a total of 5,370,356 variants were imputed and passed quality control. In SPIROMICS, a total of 5,421,262 variants were imputed and passed quality control in NHW and AA participants. Statistical analyses were performed in PLINK v1.90b3.28 and R v foss/2016b.29 Discovery analyses were performed using data from AA COPD and NHW COPD participants from COPDGene, and results were assessed in the remaining cohorts for replication. Association with weight loss for each SNP with a minor allele frequency of 5% or greater was tested assuming an additive model adjusting for age, sex, and principal components of genetic ancestry controlling for genetic ancestry. The level of genome-wide significance (GWS) for SNPs association tests was defined as $P < 5.0 \times 10^{-8}$. This level of GWS is a value traditionally used in GWAS to account for the large number of variants in linkage disequilibrium.30 It approximates a Bonferroni-corrected $P = 0.05$ for one million independent tests in the genome. Regional association plots were generated using LocusZoom31 and linkage disequilibrium information from the 1000 Genomes African Ancestry and 1000 Genomes European Ancestry reference panels32 were used. Meta-analyses were performed using the METAL33 software. Gene-based analyses were performed using the MAGMA34 software that integrates both single SNP tests with linkage disequilibrium patterns within gene regions. MAGMA assigns SNPs to genes based on physical position (±50 kb) of known genes in the NCBI site. Statistical significance (GWS) for gene-based tests was defined as $P < 2.5 \times 10^{-8}$, which corresponds to Bonferroni-corrected $P$ value threshold for $P = 0.05$ for the approximate 20 000 genes in the genome.

**Fine-mapping analysis**

To prioritize biological causality of the genotyped variants, fine mapping was performed using PAINTOR.35 PAINTOR implements a Bayesian approach incorporating genetic association results, linkage disequilibrium, and functional annotation to generate the posterior probability (PP) of causality for each variant. Fine-mapping regions were prioritized based on examination of the regional association plots as well as including variants ±25 kb from the lead variant. Single SNP test statistic ($Z$ score) information from the gene regions for the variants in the partitioned gene regions was used as input and was functionally annotated to the regions in skeletal muscle, lung, brain, adipose, and liver. Top 5 regions functionally annotated to the variants based on highest likelihood ratio were included for the analysis.
Defining modules by integrating genome-wide association study findings with protein–protein interaction data

R library dmGWAS Version 2.4 was used to integrate GWAS findings with PPI data. The dmGWAS algorithm is applied to integrate GWAS results with PPI data by using the PPI data as a search space to examine gene networks also termed modules. Nodes in the network represent gene-based results with edges representing PPI between proteins encoded by two genes. Every gene in the GWAS results is considered as an initial seed gene by the dmGWAS algorithm with a starting test statistic corresponding to the gene-based result. Using the search space defined by the PPI network additional genes are considered for inclusion in the module. A gene is included in the module if it increases the module test statistic by a factor of r. PPI data were downloaded from PINA, which includes collected and annotated data from six public databases: MINT, IntAct, DIP, BioGRID, HPRD, and MIPS/IMPact, on 16 December 2019. We further subset the PPI network data based on proteins in the Compiled Skeletal Muscle Proteome. In the current analyses, gene-based meta-analysis in the AA and NHW COPD participants was used as an input to dmGWAS. A distance constraint of d = 2 and r = 0.1 were used. A normalized module score accounting for the number of genes in the modules was generated. dmGWAS function simpleChoose was used to choose the top 10 ranked modules based on the normalized module score. Subnetworks created for each ethnicity-based PPI network was visualized using Cytoscape 3.6.1.

Pathway and tissue enrichment analyses

Gene-set enrichment analysis (GSEA) and tissue enrichment analyses was performed using the Functional Mapping and Annotation software to examine known biology of the network modules that were generated by dmGWAS. Tissue enrichment was assessed by testing whether collections of genes exhibit tissue specific expression patterns based on the Genotype-Tissue Expression project Version 8 data implemented in Functional Mapping and Annotation.

Results

Chronic obstructive pulmonary disease population characteristics

We contrasted descriptive characteristics of COPD participants in the discovery cohort, COPDGene, with the replication cohorts, SPIROMICS and ECLIPSE (Table 1). When comparing SPIROMICS and COPDGene, SPIROMICS AA COPD participants, on average, were more likely to be older, have lower BMI, have better lung function, and to have unintentional weight loss (Table 1). On average, among NHW COPD participants, those in ECLIPSE were more likely to be male participants, have lower BMI, have less smoking exposure, have worse lung function, and have higher unintentional weight loss than those in COPDGene and SPIROMICS (Table 1), whereas NHW SPIROMICS subjects tended to be older, on average, compared with COPDGene and ECLIPSE (Table 1). The prevalence of weight loss was 17% and 14.6% in COPDGene AA and NHW participants with COPD, respectively, whereas the prevalence of the weight loss trait ranged from 30.7% to 38.6% in SPIROMICS and ECLIPSE COPD participants.

Examining association of single SNPs with weight loss in participants with COPD

Among AA COPD participants, the rs35368512 variant was significantly associated with weight loss in the discovery

Table 1 Characteristics of COPD participants included in GWAS of weight loss

<table>
<thead>
<tr>
<th>Characteristic, N (%)</th>
<th>COPDGene</th>
<th>SPIROMICS</th>
<th>P value*</th>
<th>COPDGene</th>
<th>ECLIPSE</th>
<th>SPIROMICS</th>
<th>P value†</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>401</td>
<td>138</td>
<td>—</td>
<td>1380</td>
<td>1569</td>
<td>820</td>
<td>—</td>
</tr>
<tr>
<td>Sex (% male)</td>
<td>197 (49.1)</td>
<td>71 (51.4)</td>
<td>0.82</td>
<td>758 (54.9)</td>
<td>1054 (67.2)</td>
<td>466 (56.8)</td>
<td>0.48</td>
</tr>
<tr>
<td>Age</td>
<td>58.1 ± 7.5</td>
<td>61.4 ± 8.2</td>
<td>&lt;0.0001</td>
<td>64.1 ± 7.9</td>
<td>63.7 ± 7.0</td>
<td>65.9 ± 7.4</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>BMI</td>
<td>28.4 ± 6.6</td>
<td>26.7 ± 5.8</td>
<td>0.0076</td>
<td>28.6 ± 5.9</td>
<td>26.8 ± 5.6</td>
<td>27.5 ± 5.2</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Pack years</td>
<td>41.6 ± 22.4</td>
<td>42.8 ± 19.5</td>
<td>0.56</td>
<td>53.7 ± 26.1</td>
<td>50.0 ± 27.1</td>
<td>57.2 ± 51.0</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>FEV1 pp</td>
<td>56.4 ± 15.2</td>
<td>53.3 ± 16.6</td>
<td>0.046</td>
<td>54.2 ± 16.2</td>
<td>48.3 ± 15.6</td>
<td>54.1 ± 16.7</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>Weight loss a</td>
<td>68 (17.0)</td>
<td>50 (36.2)</td>
<td>0.0082</td>
<td>201 (14.6)</td>
<td>605 (38.6)†</td>
<td>252 (30.7)</td>
<td>0.0047</td>
</tr>
</tbody>
</table>

Continuous variables (age, BMI, FEV1pp, and pack-years of smoking) are represented by means and standard deviations. BMI, body mass index; COPD, chronic obstructive pulmonary disease; FEV1pp, forced expiratory volume in 1 s expressed as percentage of predicted; GOLD, Global Initiative for Chronic Obstructive Lung Disease; WL, weight loss.

*Weight loss is defined as WL > 5% and/or low BMI at any time point in the study.

†P value generated from χ2 test statistic for categorical variables and paired test statistic for continuous variables comparing African Americans between COPDGene and SPIROMICS.

P value generated from χ2 test statistic for categorical variables and one-way ANOVA test statistic for continuous variables comparing Non-Hispanic Whites between COPDGene, ECLIPSE, and SPIROMICS.
analysis [odds ratio (OR) = 3.6, 95% confidence interval (CI) = 2.3–5.6, \(P = 3.2 \times 10^{-8}\), Table S1] but did not replicate in the AA COPD participants from SPIROMICS. The rs35368512 variant is intergenic with the closest gene, \textit{GRXCR1}, located within 200 kb (Figure 1A). When the meta-analysis results of weight loss in AA COPD participants were examined, no additional variant was associated at a level of GWS (Table S2). The top variant associated with weight loss in AA COPD participants in the meta-analysis was intronic to the \textit{TBX1S} gene (Table S2). Among the NHW COPD participants in COPDGene, no single variant was significantly associated with weight loss (Table S3) nor reached GWS in the meta-analysis of all the populations (Table S4). The top single variant (rs62015138, OR = 2.1, 95% CI = 1.6–2.8, \(P = 6.4 \times 10^{-7}\)) associated with weight loss among NHW COPD participants in COPDGene was within the \textit{RBFOX1} gene (Table S3). The top single variant associated with weight loss in the meta-analysis of NHW COPD participants (rs35017521, OR = 1.4, 95% CI = 1.2–1.6, \(P = 7.7 \times 10^{-7}\), Table S4) is intergenic between a microRNA gene (\textit{MIR6072}) and a long intergenic non-protein coding RNA gene (\textit{LINC00702}).

\textbf{EFNA2 and BAIAP2 gene-based regions associated with weight loss in participants with COPD}

At the gene level, \textit{EFNA2} was associated at level of GWS with weight loss among AA COPD participants (Table 2). This finding replicated (\(P < 0.05\)) among AA COPD participants from SPIROMICS contributing to a meta-analysis \(P = 1.4 \times 10^{-8}\) (Table 2). The lead \textit{EFNA2} variant, chr19:1304013, was associated with an increased risk of weight loss (OR = 3.6, 95% CI = 1.9–6.7, \(P = 6.2 \times 10^{-5}\)) in discovery analyses, which was not GWS. In the meta-analysis, an additional two genes, \textit{C19orf24} and \textit{CIRBP}, were associated with weight loss at a level of GWS among AA COPD participants (Table 2). The \textit{EFNA2} and \textit{CIRBP} genes are located in the same region on chromosome 19 (Figure 1B). Fine mapping of 109 variants indicated 41% were needed to obtain a credible set with modest PP. Fine-mapping analyses indicated several variants within \textit{EFNA2} had modest PP of being causal (PP > 0.15, Figure 2) with the combined region accounting for 99% of the PP.

Among NHW COPD participants, the \textit{BAIAP2} gene was significantly associated with weight loss (Table 3) with the finding replicating in ECLIPSE. Among NHW COPD participants from SPIROMICS, the \textit{BAIAP2} gene was also associated with weight loss at a level near nominal significance (\(P = 0.055\)) contributing to the significant meta-analysis result (\(P = 5.20 \times 10^{-7}\)). In the discovery analyses, NHW COPD participants in COPDGene the top \textit{BAIAP2} variant, chr17:79084367, was associated with decreased risk of weight loss in COPD (OR = 0.60, 95% CI = 0.48–0.76, \(P = 1.5 \times 10^{-5}\)) but not a level of GWS. The \textit{BAIAP2} gene, on chromosome 17, is near (5 kb) to the AATK gene (Figure 3). The AATK gene is physically close but transcribed in the opposite direction with the two genes having overlapping 3’ non-coding regions.\(^{43}\) The AATK gene was significantly associated with weight loss in COPD participants from COPDGene but did not replicate in ECLIPSE or SPIROMICS (Table 3). Fine mapping of 477 variants indicated 41% were needed to obtain a credible set with modest posterior probability. Fine-mapping analysis indicated only one variant within \textit{BAIAP2} had a modest PP of being causal with the remaining variants having low likelihood of being causal (PP < 5%, Figure 3). However, the collective set which included variants within AATK increased the combined likelihood to 99%.

\textbf{Integrating GWAS findings with PPI data to identify networks of COPD weight loss genes}

Chronic obstructive pulmonary disease weight loss gene-based networks were generated using unsupervised integration of gene-based meta-analysis results with PPI data of proteins expressed in skeletal muscle. Integration of the meta-analysis weight loss gene-based results from AA COPD participants identified 12,135 modules. A consensus module was generated from the top 10 most significant modules included 29 genes (Figure 4, Table S5). Several of these genes (\textit{EFNA2}, \textit{CIRBP}, \textit{WDR88}, and \textit{KCNJ3}) in the consensus module were among the top 10 most associated with weight loss in AA COPD participants (Table 2). GSEA indicated the consensus module genes associated with weight loss in the AA COPD participants were enriched in pathways involved in \textit{NRF1} signalling, adipogenesis, synapse, and RNA metabolism among others (Table S6). Integration of the meta-analysis weight loss gene-based results from NHW COPD participants with PPI network data identified 12,168 modules. A consensus network was created from the top 10 most significant modules and was comprised of 36 genes (Figure 5, Table S7). Of which, several genes (\textit{BAIAP2}, \textit{AATK}, \textit{ZZEF1}, and \textit{RHOB}) were among the top 10 most significantly associated with weight loss in NHW COPD participants (Table 3). GSEA results indicated the consensus module genes were enriched in pathways involved in adipogenesis, synapse signalling, Rho GTPase signalling as well as genes in other known other pathways (Table S8). Despite having only one gene, \textit{UBC}, common to both the AA and NHW weight loss consensus networks, there were six known gene-sets in common. These were comprised of genes (i) involved in synapse signalling; (ii) involved in formation of the incision complex; (ii) with sites recognized by miR-520D; (iv) involved in protein tagging for modification, sequestration, transport, or degradation; (v) involved in adipocyte differentiation; and (vi) involved in subdivision of chromosomal regions.

The \textit{EFNA2} gene is a member of the set of 16 synapse signalling genes represented in consensus weight loss networks in both AA and NHW COPD participants (combined GO
**Figure 1** (A) Regional association plot for top weight loss variant region in African American participants from COPDGene near variant, 4:43232667. (B) Regional association plot for top weight loss variant region in non-Hispanic White COPD participants from COPDGene on chromosome 19. In both panels, lead SNPs denoted in purple. Colour of remaining SNPs indicates degree of linkage disequilibrium with the lead SNP, as measured by $r^2$, the squared coefficient of correlation.
Table 2 Based on meta-analysis, top 10 genes associated with weight loss in participants with COPD in African American cohorts (COPDGene and SPIROMICS)

<table>
<thead>
<tr>
<th>Gene</th>
<th>CHR</th>
<th>START</th>
<th>STOP</th>
<th>NSNPS</th>
<th>ZSTAT</th>
<th>P value</th>
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<th>ZSTAT</th>
<th>P value</th>
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<tbody>
<tr>
<td>EFNA2</td>
<td>19</td>
<td>1236153</td>
<td>1351430</td>
<td>252</td>
<td>5.6</td>
<td>1.40E-08</td>
<td>270</td>
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<td>C19orf24</td>
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<td>81</td>
<td>4.6</td>
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COPD, chronic obstructive pulmonary disease.

*Denotes genes that appear in the consensus network.

Figure 2 Fine mapping of EFNA2 region associated with weight loss among African American COPD participants. (A) Scatterplot of location vs. posterior probabilities with credible set, (B) physical position of EFNA2, and (C) functional annotation tracks.
SYNAPSE gene list in Tables S6 and S8). The 16 genes include STXBP3, DISC1, KCNK1, KPNA2, ITSN1, and PPP2CA as well as PPP1CA, SNAP23, NSF, ELAVL1, BIN1, PACSIN2, ARL8B, STX7, and STXBPS in the AA and NHW COPD weight loss consensus networks, respectively. Tissue specificity analyses indicated genes were significantly down-regulated in the liver ($P < 1 \times 10^{-3}$). Further, five of these genes (KCNK1, ELAVL1, ARL8B, PPP2CA, and STXBP5) also have sites recognized by the miRNAS20D (GSEA adjusted $P = 2.1 \times 10^{-3}$).

**Discussion**

In the current report, we identified two genes significantly associated with weight loss in COPD: EFNA2 and BAIAP2. Although, EFNA2 and BAIAP2 were associated with weight loss in COPD among AA and NHW participants, respectively, they were also members of the COPD weight loss consensus networks. This indicates there is genetic variation in these genes that could influence other genes within the same network. As COPD and weight loss are complex traits, we expect genetic variation in many genes contribute to dysregulation at the pathway level. Importantly, many of the gene-sets enriched in the COPD weight loss consensus networks have important roles in skeletal muscle regeneration and tissue remodelling.

The EFNA2 gene encodes the membrane-bound protein ephrin-A2. Ephrins interact with Eph receptors via contact-dependent cell–cell signalling regulating developmental processes and adult tissue homeostasis.44 Interestingly, ephrin-A2 participates in bidirectional signalling by activating eph receptors on neighbouring cells as well as its own downstream pathways and has been shown to negatively regulate progenitor cell proliferation.45 The EFNA2 gene is a member of the GO SYNAPSE gene-set whose members were represented in both the AA and NHW COPD weight loss consensus networks of genes. We demonstrated using Genotype-Tissue Expression data the 15 COPD weight loss genes in the SYNAPSE gene-set are down-regulated in the liver. The role in the liver is interesting as in metabolic disturbances have been shown to start in the liver before progressing to adipose and skeletal muscle tissues in mouse cancer cachexia models.46,47 Further, the liver has afferent and efferent neurons which may influence appetite and hormone signals.48 However, we were not able to test whether genetic variation associated with weight loss in COPD leads to altered gene expression of these 15 genes in the liver. Therefore, additional research into whether expression of these genes is up-regulated in the liver with COPD cachexia and weight loss is needed.

The BAIAP2 gene encodes the insulin-responsive protein of mass 53 kD (IRSp53), which is an adaptor protein primarily known for its role in modulating actin dynamics and membrane protrusions in cell to cell signalling.49 Impaired skeletal

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**Table 3** Top 10 genes associated with weight loss in participants with COPD based on meta-analysis in non-Hispanic White cohorts (COPDGene, ECLIPSE, and SPIROMICS)

<table>
<thead>
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<th>NSNPS</th>
<th>ZSTAT</th>
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COPD, chronic obstructive pulmonary disease; Denotes genes that appear in the consensus network.
muscle regeneration is one mechanism contributing to skeletal muscle loss in cachexia. IRSp53 can act as a negative regulator of myogenic differentiation influencing the development of skeletal muscle as well skeletal muscle regeneration. Previous research demonstrated COPD patients may exhibit heterogeneous and distinct skeletal muscle molecular biomarker patterns in response to pulmonary rehabilitation. However, IRSp53 was not among the biomarkers investigated in the previous research. It is possible dysregulation of IRSp53 may be one mechanism contributing to impaired ability to regenerate skeletal muscle in cachexia; however, this requires further research. Also, GSEA analyses highlighted BAIAP2 involvement in Rho signalling. Activation of the Rho signalling pathway is required for the maintenance of myotubes. Interestingly, ephrins such as ephrin-B whose gene were associated with weight loss among AA COPD participants activate eph receptors who exert downstream by regulating Rho GTPase signalling.

Our study has many strengths but also limitations. Strengths of the study include a large sample size of COPD participants in multiple cohorts followed longitudinally with weight loss and genotype data available. We also used innovative network methods, going beyond generating a list of genes associated with weight loss in COPD, providing

Figure 3  Fine mapping of BAIAP2 region associated with weight loss among non-Hispanic White COPD participants. (A) Scatterplot of location vs. posterior probabilities with credible set, (B) physical position of genes including BAIAP2 in region, and (C) functional annotation tracks.
rationales for further mechanistic research. Limitations of the study include heterogeneity in number of visits used to define weight loss in the discovery and replication cohorts. In COPDGene, a self-reported unintentional weight loss greater than 5% in the past year or low BMI collected at a single visit was used. Whereas in ECLIPSE and SPIROMICS, weight loss and BMI were measured at several annual visits. The prevalence of weight loss was higher in ECLIPSE and SPIROMICS which may be due to increased opportunities for observing weight loss over the course of the study. In COPDGene, weight loss was also unintentional whereas in the other two studies participants may have been intentionally trying to lose weight which could inflate the prevalence of weight loss. Self-reported, unintentional weight loss is likely a more conservative measure, however, may also be subject to recall bias. Discovery analyses were performed using more strict criteria to code the weight loss trait which would have biased findings towards the null. However, the lower number of visits over larger time intervals in COPDGene likely led to misclassification and loss of follow-up of participants with COPD who passed away before weight loss could be recorded by the study. This would have limited our ability to identify some true associations with weight loss in COPD using COPDGene as discovery. Future studies of weight loss in COPD should aim to collect weight measurements more frequently such as the intervals employed in ECLIPSE and SPIROMICS. Further, we also employed a Bonferroni-corrected level of significance to gene-based findings which may have been overly conservative.

Furthermore, the UBC gene encoding Ubiquitin C was the only gene in common between the AA and NHW COPD weight loss consensus networks. The ubiquitin-proteasome system is fundamental to muscle atrophy in cachexia. However, we previously demonstrated the network analysis method, dmGWAS, was sensitive to hub genes such as UBC, encoding Ubiquitin C. For these reasons, we also performed the network analyses excluding UBC and found EFNA2 and BAIPA2 were robustly included in each consensus module whether UBC was included in the PPI search space or not. Finally, we analysed genotyped and imputed SNP data, which led to the identification of two gene regions associated with weight loss in COPD rather than specific genetic variants. We maximized the information in the regions through our fine-mapping approach. Although we analysed a combined

Figure 4 COPD weight loss consensus network generated from African American participant analyses. Node size is proportional to P value significance where the bigger the node size the smaller the P value from the gene-based meta-analysis result. Nodes (circles) represent genes that are among the top genes in the consensus network. Edges (lines) are known protein–protein (PPI) interactions. Nodes filled in black represent genes robust to exclusion of UBC from the PPI network.
set of 4308 subject with COPD in the three studies combined, the samples sizes became small when stratified by study and ancestry group likely limiting the power to replicate findings in the discovery. For example, the top variant associated with weight loss in the meta-analysis of AA COPD subjects is intronic to the \textit{TBX15} gene. \textit{TBX15} is a member of the T-box family of transcription factors and has been previously associated with waist to hip to ratio.\textsuperscript{55} Nonetheless, we were able to maximize information using our integrative approach to discover new aetiology for weight loss in COPD. However, an expanded analysis in these populations of COPD participants using whole-genome sequence with integration with other omics data may lead to data identifying specific genetic variants, which may guide personalize medicine approaches.

To summarize, \textit{BAIAP2} and \textit{EFNA2} genes were significantly associated with weight loss in COPD among NHW and AA participants. Our integrative network analyses identified COPD weight loss genes enriched with genes involved in skeletal muscle regeneration and tissue remodelling as well as providing rationales for further mechanistic research. Identification of genetic variation contributing to weight loss in COPD due to impaired skeletal muscle regeneration and tissue remodelling may enable discovery of therapies that could enhance response to pulmonary rehabilitation.

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SPIROMICS

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Conflict of interest

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Online supplementary material

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1. COPD GOLD Stage ≥ 2 participants with phenotype and genotype data included in analyses. A) COPD participants included from the COPDGene study; B) COPD participants included from SPIROMICS study; C) COPD participants included from ECLIPSE study. GWAS – Genome Wide Association Study. In COPDGene, 2 subjects with Visit 2 self-reported weight loss data were excluded for being incongruent with Visit 1 weight.

Table S1. SNPs associated with COPD weight loss among African American COPD participants from COPDGene (N = 401) with P < 1E-5.

Table S2. Top 10 SNPs associated with weight loss among African American COPD participants based on meta-analysis COPDGene and SPIROMICS.

Table S3. SNPs associated with COPD weight loss among Non-Hispanic White COPD participants from COPDGene with P < 1E-5.

Table S4. Top 10 SNPs associated with weight loss among Non-Hispanic White COPD participants based on meta-analysis of three studies (ECLIPSE, COPDGene and SPIROMICS).

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Table S5. Genes represented in the COPD weight loss consensus network from African American participants with corresponding gene-based results provided.

Table S6. Gene Set Enrichment Analysis of Consensus Network Genes associated with weight loss among AA COPD participants from COPDGene and SPIROMICS.

Table S7. Genes represented in the COPD weight loss consensus network from non-Hispanic White participants with corresponding gene-based results provided.

Table S8. Gene Set Enrichment Analysis of Consensus Network Genes associated with weight loss among NHW COPD participants from COPDGene, SPIROMICS and ECLIPSE.

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