Human CCL3L1 copy number variation, gene expression, and the role of the CCL3L1-CCR5 axis in lung function [version 1; referees: 1 approved]

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Abstract

Background: The CCL3L1-CCR5 signaling axis is important in a number of inflammatory responses, including macrophage function, and T-cell-dependent immune responses. Small molecule CCR5 antagonists exist, including the approved antiretroviral drug maraviroc, and therapeutic monoclonal antibodies are in development. Repositioning of drugs and targets into new disease areas can accelerate the availability of new therapies and substantially reduce costs. As it has been shown that drug targets with genetic evidence supporting their involvement in the disease are more likely to be successful in clinical development, using genetic association studies to identify new target repurposing opportunities could be fruitful. Here we investigate the potential of perturbation of the CCL3L1-CCR5 axis as treatment for respiratory disease. Europeans typically carry between 0 and 5 copies of CCL3L1 and this multi-allelic variation is not detected by widely used genome-wide single nucleotide polymorphism studies.

Methods: We directly measured the complex structural variation of CCL3L1 using the Paralogue Ratio Test and imputed (with validation) CCR5del32 genotypes in 5,000 individuals from UK Biobank, selected from the extremes of the lung function distribution, and analysed DNA and RNAseq data for CCL3L1 from the 1000 Genomes Project.

Results: We confirmed the gene dosage effect of CCL3L1 copy number on CCL3L1 mRNA expression levels. We found no evidence for association of CCL3L1 copy number or CCR5del32 genotype with lung function.

Conclusions: These results suggest that repositioning CCR5 antagonists is unlikely to be successful for the treatment of airflow obstruction.
Introduction

Genome-wide association studies have identified thousands of disease-gene associations leading to new disease insight and potential new approaches to treatment. It has been shown that drug targets supported by genetic studies have an increased chance of success in clinical development. Even so, only a subset of candidate drugs will make it through to the clinic. Identifying opportunities for repositioning existing drugs and targets is therefore an appealing prospect and using genetic studies to define alternative indications for an already-approved drug is a promising approach.

The MIP-1alpha (encoded by \textit{CCL3} and \textit{CCL3L1})-CCR5 signaling axis is important in a number of inflammatory responses, including macrophage function, and T-cell-dependent immune responses. It is perturbed by CCR5 antagonists such as Pfizer’s maraviroc, the only CCR5 antagonist to be approved by the United States Food and Drug Administration. Identification of a genetic association of variants within the genes involved (\textit{CCR5} and \textit{CCL3L1}) would strongly support the potential use of CCR5 antagonists in the treatment of respiratory conditions.

In mice, MIP-1alpha is implicated in virus-mediated inflammation of the lung, pulmonary eosinophilia following paramyxovirus infection, clearance of pulmonary infections, and in the response to respiratory syncytial virus infection. In humans, Mip-1alpha controls the recruitment of immune cells to inflammatory foci, and increased levels of MIP-1alpha mRNA are found in bronchial epithelial cells of COPD patients, and increased protein levels in the sputum of COPD patients where increased macrophage and neutrophil infiltration in the lung is a key pathology.

The \textit{CCR5} gene in humans has a 32bp exonic deletion allele (rs433, \textit{CCR5d32}) with a minor allele frequency of between 5–15% in Europeans. This allele causes a translational frameshift and abolishes expression of the receptor at the cell surface, such that homozygotes for the deletion allele lack any functional CCR5 receptor. This variant has been strongly and repeatedly associated with resistance to HIV infection and slower HIV progression, as CCR5 is a common coreceptor for HIV entry into T-lymphocytes. The \textit{CCR5d32} allele has been suggested to confer a reduced risk of asthma in children in one study although this has not been replicated.

In humans, there are two isoforms of MIP-1alpha, the LD78a isoform encoded by the \textit{CCL3} gene and the LD78b isoform encoded by the paralogous \textit{CCL3L1} gene. The two isoforms differ by three amino acids, but only one of these small changes, a serine to proline change at position 2 of the mature protein, alters the affinity to the cell surface receptor CCR5, with the beta isoform (\textit{CCL3L1}) having approximately six-fold greater affinity for CCR5 than the alpha isoform (\textit{CCL3}).

The \textit{CCL3L1} gene is part of a complex structurally variable region, although the \textit{CCL3} gene is not. The \textit{CCL3L1} gene and the neighboring \textit{CCL4L1} gene are tandemly repeated with the total diploid copy number ranging from 0 copies to 6 copies in Europeans. Higher copy numbers are observed elsewhere, for example 10 in Tanzanian and 14 in Ethiopian populations. Previous studies have shown evidence of a gene dosage effect, with \textit{CCL3L1} gene dose reflected in mRNA levels as well as in the ability to chemotact monocytes.

Measuring \textit{CCL3L1} multiallelic copy number variation has been challenging. Early studies used qPCR assays with a low signal-to-noise ratio, but assays based on the paralogue ratio test (PRT), allowed more accurate estimation of diploid copy number. Because of the challenges in measuring \textit{CCL3L1} copy number in sufficiently large and well-powered sample sizes, the effect of structural variation of the genes encoding the MIP-1alpha-CCR5 ligand-receptor pair has not been adequately explored.

In this study, we set out to confirm previous reports that \textit{CCL3L1} copy number is associated with \textit{CCL3L1} gene expression, then measure \textit{CCL3L1} copy number and \textit{CCR5d32} genotype in 5000 individuals from UK Biobank, and finally test for association with lung function. Furthermore, we validated our copy number typing approach and observed copy number frequencies using publicly available sequence data from the 1000 Genomes Project. For \textit{CCL3L1} copy number measurement in the 5000 individuals from UK Biobank, we used a triplex PRT, which is considered to be the gold standard approach for measurement of this copy number variation. For genotyping of \textit{CCR5d32} in UK Biobank, we used a standard genotype imputation approach with additional PCR validation. We tested for association with extremes of Forced Expired Volume in 1 second (FEV \textsubscript{1}) as a binary trait. This study is the largest analysis of the effect of \textit{CCL3L1} copy number and \textit{CCR5d32} genotypes on lung function undertaken to date.

Methods

Sample selection

Individuals were selected from the UK BiLEVE subset of UK Biobank. Data from the UK BiLEVE study are available at http://www.ukbiobank.ac.uk/data-showcase/. In brief, 502,682 individuals were recruited to UK Biobank of whom 275,939 were of self-reported European-ancestry, and had two or more measures of Forced Expired Volume in 1s (FEV \textsubscript{1}) and Forced Vital Capacity (FVC) measures (Vitalograph Pneumotrac 6800, Buckingham, UK) passing ATS/ERS criteria. Based on the highest available FEV \textsubscript{1} measurement, 50,008 individuals with extreme low (n=10,002), near-average (n=10,000) and extreme high (n=5,002) % predicted FEV \textsubscript{1} were selected from amongst never-smokers (total n=105,272) and heavy-smokers (mean 35 pack-years of smoking, total n=46,758), separately. For this study, we selected 2500 age-matched European-ancestry heavy smokers from the extreme high and extreme low % predicted FEV \textsubscript{1} subsets defined for the UK BiLEVE study (Figure 1, Table 1). DNA samples for these 5000 individuals were prepared by UK Biobank and provided back to the University of Leicester with new identification codes such that typing of \textit{CCL3L1} copy number and \textit{CCR5d32} was blinded to lung function status. Positive control samples for the copy number typing were...
**Figure 1. Study design.** FEV₁ is percent predicted FEV₁. *Lung function measurement quality control defined previously*.[33] *Final numbers after quality control*.[33]

**Table 1. Demographics of selected UK Biobank cohort.**

<table>
<thead>
<tr>
<th></th>
<th>Low FEV₁ (n=2500)</th>
<th>High FEV₁ (n=2500)</th>
</tr>
</thead>
<tbody>
<tr>
<td>n (%) male</td>
<td>1250 (50%)</td>
<td>1250 (50%)</td>
</tr>
<tr>
<td>Age</td>
<td>56.9 / 7.9 (40, 70)</td>
<td>56.9 / 7.9 (40, 70)</td>
</tr>
<tr>
<td>Pack-years</td>
<td>40.6 / 22.5 (10.8, 301.0)</td>
<td>29.37 / 13.4 (10.5, 134.0)</td>
</tr>
<tr>
<td>Pack-years as a proportion of lifespan</td>
<td>0.96 / 0.47 (0.42, 7.00)</td>
<td>0.70 / 0.29 (0.42, 3.03)</td>
</tr>
<tr>
<td>FEV₁ (litres)</td>
<td>1.50 / 0.47 (0.36, 3.38)</td>
<td>3.64 / 0.73 (2.02, 6.72)</td>
</tr>
<tr>
<td>Percent predicted FEV₁</td>
<td>51.4 / 11.0 (14.9, 74.5)</td>
<td>123.3 / 8.2 (112.8, 205.7)</td>
</tr>
</tbody>
</table>

Values are Mean / SD (range), unless stated.

from the Human Random Control panel from Public Health England (C0075 – 1 copy, C0150 – 2 copies, C0007 – 3 copies, C0877 – 4 copies), as used previously.[26]

CCL3L1 copy number estimation in UK Biobank and 1000 Genomes Project samples using the paralogue ratio test (PRT)

*CCL3L1* copy number was determined using a triplex paralogue ratio test (PRT) assay as used previously.[24,26]. Briefly, PRT is a comparative PCR method that amplifies a test and reference locus using the same pair of primers, followed by capillary electrophoresis and quantification of the two products.[32,36]. The triplex assay produced three independent estimates of copy number per test, of which the average was taken as a representative copy number value. The three values were consistent in 95% of samples, however, for 5% of samples the value from the LTR61A PRT assay was significantly lower than the other two PRT values, and an average of the two consistent PRTs was taken in these 5% of samples. For each typing experiment, 4 positive controls of known copy number were also included, as previously.[26,37]. The copy number values clustered about integer copy numbers, and a Gaussian mixture model was fitted to allow assignment of individuals to an integer copy number call using CNVtools.[38]. For the 5000 individuals from UK Biobank, 58 individuals were selected by UK Biobank investigators as blind spiked duplicates as part of the quality control check to ensure genotyping accuracy. Copy numbers from UK Biobank samples are available from UK Biobank at [http://www.ukbiobank.ac.uk/data-showcase/](http://www.ukbiobank.ac.uk/data-showcase/).

Gene expression levels in 1000 genomes project lymphoblastoid cell lines

Matched RNAseq data that is publically available for the 1000 genomes samples were grouped based on *CCL3L1* copy number and analysed for their differential expression using Cufflinks v2.1.1.[39]. This allows measurement of the effect of genomic copy number of *CCL3L1* on gene expression levels. The analyses were all performed on ALICE High Performance Computing Facility at the University of Leicester. The RNAseq data were downloaded from EBI ArrayExpress (accessions E-GEUV-1, E-GEUV-2, E-GEUV-3).[40]. Using Cufflinks, the fragments per
kilobase of transcript per million fragments mapped (FPKM) values were estimated by applying a statistical model that normalises the mapped reads by length and their abundance. Briefly, the fragment reads are divided by transcript size and the total number of reads and then adjusted to 1 kb and 1 million reads.

Genotyping of CCR5d32 polymorphism

Imputation to 1000 Genomes Project Phase 1+UK10K reference panel^{28} and PCR were used to genotype the CCR5d32 polymorphism (rs333) in the 5000 UK Biobank individuals. Phasing and imputation were undertaken with SHAPEIT v2.r790^{32} and IMPUTE2 v2.3.1^{41}. For individuals with imputation posterior probability <0.95 (431 samples), and an additional 20 samples that were imputed as homozygous for the minor del32 allele, we validated the imputation results using direct PCR genotyping. Duplicates of a random selection of 28 of individuals were included as a quality control check for genotyping reproducibility (genotyping was also blinded to duplicate status). Genotypes from UK Biobank samples are available from UK Biobank at http://www.ukbiobank.ac.uk/data-showcase/.

CCL3L1 copy number estimation from sequencing data for 1000 Genomes Project individuals

1000 genomes phase 3 whole genome aligned Bam files generated from Illumina platforms available from the European Bioinformatics Institute were validated and the genomic region including CCL3L1 (hg19:chr17:33670000-35670000) was analysed using CNVrd2^{44}. Using 500bp window sequence read depth, the sequence read depth was calculated across the region for all 2502 genomes from 26 populations, and standard deviation/quantile calculated for each window. The segmentation scores obtained from this analysis were clustered into different groups using a Gaussian mixture model. A prior information for all populations was estimated using the expectation maximisation (EM) algorithm on a population group with clear clusters of segmentation scores. The prior information (means, standard deviations and proportions of the mixture components) was fed into Bayesian model to infer CCL3L1 integer copy number in all populations. Copy number estimates are available from dbVar under study accession number nstd155.

Association analysis

We tested for association of CCL3L1 copy number and CCR5d32 genotype separately with lung function extremes (as a binary trait) using logistic regression in R v. 3.2.3 with pack-years of smoking and the first ten principal components (obtained previously using full genome-wide SNP genotyping data) to adjust for fine-scale population structure as covariates^{39}. For CCR5d32, a genotypic genetic model was assumed for the primary analysis. We then fitted a full linear regression model that included CCR5d32 genotype (genotypic mode), CCL3L1 copy number, pack years, 10 principal components and a term for the interaction of CCR5d32 and CCL3L1.

A previous version of this manuscript is available on Biorxiv https://www.biorxiv.org/content/early/2018/01/17/249508

Results

Using CNVrd2, we typed CCL3L1 copy number from whole genome sequence alignments for 2502 individuals from the 1000 Genomes project (Figure 2a). The data were grouped into large superpopulations, as defined by the 1000 Genomes Project^{38}, and our analysis confirmed previous observations that Europeans have the lowest CCL3L1 diploid copy number, ranging between 0 and 5 with a mean copy number of 1.97, and sub-Saharan Africans have the highest diploid copy number, ranging between 1 and 9 with a mean of 4.19, which is more than twice as high as Europeans (Table 2)^{39,42}.

For 144 individuals from the CEU (n=96) and YRI (n=48) populations of the 1000 Genomes project, we also determined CCL3L1 copy number using the PRT approach (Figure 2b). There was strong concordance between results, with discrete clusters of raw data, representing individual integer copy numbers, formed, particularly at low copy number. For the range seen in Europeans (copy numbers 0 to 5), there are seven clear discrepancies, which gives a joint error rate of 5%.

To confirm previous studies that reported an association between CCL3L1 copy number and CCL3L1 mRNA levels, we compared the 1000 Genomes Project CCL3L1 copy numbers with transcript levels of CCL3L1 and its non-copy number variable parologue CCL3, as generated by RNAseq of the corresponding B-lymphoblastoid cell lines (Figures 3a and b). Comparison with transcript level estimates using RNAseq data showed a clear positive correlation between CCL3L1 copy number and expression level (Figure 3b, r=0.25, p<2x10^{-6}).

Having confirmed a relationship between gene copy number and transcript expression level (CCL3L1), we investigated the relationships between CCL3L1 copy numbers, CCR5d32 genotype and lung function in individuals selected from the extremes of the lung function distribution in UK Biobank. We typed 5000 UK Biobank samples using PRT, with 19 failures. The results showed a clear mixture of Gaussian distributions centered on each integer copy number (Figure 2c). All 58 duplicates were consistently typed, resulting in an error rate between 0% and 4.7%. We observed clear distances between the clusters, further suggesting that the measurement error rate for this cohort is likely to be low.

We estimated CCL3L1 integer copy numbers in all the samples using Gaussian mixture modelling (Table 3). The copy number
Figure 2. CCL3L1 Copy number typing. (a) Histogram of raw copy number estimates of 1000 Genomes Project samples from sequence read depth represented as segmentation scores on the x axis, generated by CNVrd2, with higher scores reflecting higher copy number. (b) Validation of 144 1000 Genomes Project samples using PRT (x axis) against estimates made from sequence read depth. Colours/symbols in the scatterplot represent different integer copy numbers inferred from PRT clusters. (c) Histogram of raw copy number estimates using PRT for the UK Biobank cohort.
Table 2. CCL3L1 copy number frequency distributions in 1000 Genomes data.

<table>
<thead>
<tr>
<th>Superpopulation</th>
<th>n</th>
<th>average copy number</th>
<th>minimum copy number</th>
<th>maximum copy number</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFR (Sub-Saharan African)</td>
<td>661</td>
<td>4.19</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>AMR (Admixed American)</td>
<td>347</td>
<td>2.71</td>
<td>0</td>
<td>8</td>
</tr>
<tr>
<td>EAS (East Asian)</td>
<td>504</td>
<td>3.52</td>
<td>0</td>
<td>9</td>
</tr>
<tr>
<td>EUR (European)</td>
<td>501</td>
<td>1.97</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>SAS (South Asian)</td>
<td>489</td>
<td>2.39</td>
<td>0</td>
<td>7</td>
</tr>
</tbody>
</table>

Figure 3. Copy number and expression level of CCL3L1 and CCL3 in lymphoblastoid cell lines. (a) CCL3 mRNA level (FPKM units) across different CCL3L1 copy numbers. (b) CCL3L1 mRNA level (FPKM units) across different CCL3L1 copy numbers. (c) CCL3L1:CCL3 mRNA ratio across different CCL3L1 copy numbers. Individual data points are shown, with red bars indicating median and interquartile ranges.
number range was consistent with previous observations in UK population\textsuperscript{39}, and with our estimation from the 1000 Genomes project samples. The two copy genotype was the most frequent with a frequency of 0.563. The CCL3L1 zero copy null genotype is uncommon, with a frequency of 2.5\% in the UK. 4993 of the 5000 UK Biobank samples were genotyped for CCR5d32 by imputation with the genotypes for 474 individuals validated using direct PCR analysis. There was no evidence that the genotype frequencies departed from Hardy-Weinberg equilibrium (chi-squared test, p=0.35) and the observed CCR5d32 deletion allele frequency was 0.11, consistent with previous estimates\textsuperscript{41}.

A total of 4975 UK Biobank individuals had both CCL3L1 copy number and CCR5d32 genotypes measured (2486 high and 2489 low FEV\textsubscript{1}, Table 4). There was no evidence of an association between CCL3L1 copy number and CCR5d32 genotype (chi-squared test p=0.803).

We fitted a full model with both CCR5 genotypes (genotypic model) and CCL3L1 copy number and an interaction term as described above. This was undertaken in order to identify whether particular combinations of CCL3L1 copy number and CCR5d32 genotype were differentially associated with lung function. Pack years of smoking and 10 principal components were included as covariates. No associations were significant (Table 5).

**Table 3. CCL3L1 copy number counts in UK Biobank data.**

<table>
<thead>
<tr>
<th>CCL3L1 diploid copy number</th>
<th>Number of samples</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>127</td>
<td>0.025</td>
</tr>
<tr>
<td>1</td>
<td>1046</td>
<td>0.210</td>
</tr>
<tr>
<td>2</td>
<td>2806</td>
<td>0.563</td>
</tr>
<tr>
<td>3</td>
<td>853</td>
<td>0.171</td>
</tr>
<tr>
<td>4</td>
<td>128</td>
<td>0.026</td>
</tr>
<tr>
<td>5</td>
<td>21</td>
<td>0.004</td>
</tr>
<tr>
<td>Sum</td>
<td>4981</td>
<td>0.999</td>
</tr>
</tbody>
</table>

**Table 4. CCR5d32 genotype counts by CCL3L1 copy number in UK Biobank data.**

<table>
<thead>
<tr>
<th>CCL3L1 copy number</th>
<th>CCR5d32 genotype</th>
<th>OR (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ref/ref</td>
<td>del32/ref</td>
<td>0.84 (0.57-1.23)</td>
<td>0.38</td>
</tr>
<tr>
<td>del32/ del32</td>
<td></td>
<td>0.29 (0.07-1.30)</td>
<td>0.11</td>
</tr>
<tr>
<td>ref/ref</td>
<td>del32/homo</td>
<td>1.00 (0.92-1.09)</td>
<td>0.97</td>
</tr>
<tr>
<td>del32/homo</td>
<td></td>
<td>1.11 (0.93-1.32)</td>
<td>0.27</td>
</tr>
<tr>
<td>ref/ref</td>
<td>del32/interact</td>
<td>1.74 (0.83-3.64)</td>
<td>0.14</td>
</tr>
</tbody>
</table>

2486 samples with high FEV\textsubscript{1} and 2489 samples with low FEV\textsubscript{1}.

Covariates: smoking pack-years, 10 principal components of SNP genetic variation.

**Table 5. Association analysis of CCR5 genotype and CCL3L1 copy number with high vs low FEV\textsubscript{1}.**

Discussion

Our study provides robust large-scale confirmation of a gene dosage effect of CCL3L1 copy number on CCL3L1 mRNA levels, and also emphasises the strong dependence of CCL3L1:CCL3 mRNA ratio on copy number, with CCL3L1 copy number accounting for 50\% of total variation. Although it is clear that CCL3L1 is expressed at much lower levels than CCL3, the MIP-1alpha isoform encoded by CCL3L1 (LD78beta) has a much stronger affinity to the CCR5 receptor than MIP-1alpha isoform CCL3 (LD78alpha). It therefore seems likely that the CCL3L1 copy number variation mediates a biological effect in vivo. It should be noted that the expression data are from transformed B-lymphoblastoid cell lines, but a gene dosage effect is consistent with a study using fresh monocytes from 55 different individuals stimulated with bacterial lipopolysaccharide\textsuperscript{38}.

Our analysis provides evidence that there is no effect of either CCL3L1 copy number or CCR5d32 genotype, or any combinations of genotypes at the two loci, on lung function. This suggests that, although the MIP-1alpha-CCR5 signaling axis can be disrupted by artificial CCR5 antagonists, there is no evidence that this axis has a functional effect on lung function and that development of new drugs to target this axis, or repurposing of existing drugs, might be of little or no therapeutic benefit in treating COPD.

We analysed approximately 5000 individuals. Whilst this represents a large sample size for labour-intensive PRT assays, it is a modest sample size in comparison with those employed in GWAS. That said, power was boosted by selecting from the extremes of the lung function distribution in the very large (n~500,000) UK Biobank.

We reported PRT error rates of 2.5\% for the 144 1000 Genomes Project samples and between 0\% and 4.75\% for the 4981 UK Biobank participants. A previous study using this PRT approach...
estimated an error rate of less than 0.1%, which suggests that much of the joint error rate for the PRT and sequence read depth could be due to errors in the sequence read depth approach.

The exact boundaries of the CCL3L1 CNV have yet to be determined with precision but it is known to include the CCL4L1 gene, which encodes MIP-1beta. The human genome assembly GRCh38 shows a single copy CCL3L1/CCL4L1 repeat unit, and also includes the TBC1D3 gene, encoding TBC1 Domain Family Member 3. The GRCh38 alternative assembly chr17_KI270909v1_alt shows two repeat units, both including TBC1D3. However an earlier assembly shows a complete contig with two repeat units carrying CCL3L1/CCL4L1, only one of which carries TBC1D3. ArrayCGH and fiber-FISH both confirm this is real heterogeneity by showing that the TBC1D3 gene is included in some, but not all, tandemly repeated units in some individuals, together with CCL3L1 and CCL4L1. Throughout this paper, and in most of the literature, CCL3L1 CNV is used as a shorthand to describe the CNV of this complex repeat unit.

Given the gene content of this repeat unit, we would expect a gene dosage effect for CCL4L1 and TBC1D3, in addition to CCL3L1, but this has not yet been confirmed. Our data do, however, show no effect of CCL3L1 copy number on expression levels of its close parologue, CCL3, which is immediately proximal to the CNV. This difference shows that the considerable variation in genome structure distal to the CCL3 gene does not affect overall levels of CCL3 expression.

In summary, we selected individuals from the extremes of the lung function distribution of a very large general population cohort. We found no association of CCL3L1 copy number, nor of the CCR5d32 variant with lung function, as defined by FEV1.

Data availability
UK Biobank data are available upon application to the UK Biobank (https://www.ukbiobank.ac.uk/) to all bona fide researchers. Access details can be found at: http://www.ukbiobank.ac.uk/register-apply/.

Data from the UK BiLEVE study are available at http://www.ukbiobank.ac.uk/data-showcase/.

Competing interests
AM, SJ and IK are employees of Pfizer, Inc. All other authors have no competing interests.

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LVW holds a GSK / British Lung Foundation Chair in Respiratory Research.

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References

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This exploration of association between lung function and CCL3L1 copy number and CCR5 deletion status has the interesting motivation that if the CCR5 signalling axis is associated with lung function, CCR5 antagonists might be repurposed for treatment of respiratory disease. I thought that the methods adopted were well-powered, the analyses appropriate and the conclusions drawn well justified. The mix of information from different cohorts and different genetic analyses was carefully handled to allow powerful conclusions to be derived from high-throughput data sources, while at the same time ensuring that genotypes and copy numbers were properly substantiated by direct DNA typing.

CNVrd2 does a good job of extracting copy number states for CCL3L1/CCL4L1 from 1000 Genomes Project sequence data, and the concordance in directly typed samples (Figure 1) is impressive. The clustering of the results from the UK Biobank samples is also very clear, validating the precision and accuracy of the Gaussian mixture models extracted by CNVtools.

For the gene expression analyses at the RNA level I would agree with the conclusions stated in the first paragraph of the Discussion, that although the transcription level for CCL3L1 is much lower than for CCL3, the greater biological effect of each CCL3L1 protein molecule is likely to mean that the variation attributable to CNV has genuine function effect.

Overall, I think this is an interesting study carried out to a high standard of technical accuracy and robustness, and I have no suggestions for its improvement. One suggestion I offer the authors for their consideration is the possibility of using local SNPs to impute CCL3L1 CN from Biobank data. Clearly, any imputation of a multiallelic CNV from diallelic SNPs is likely to have limited power, but even incomplete imputation of CCL3L1 CN from SNP data alone may unlock power by allowing analysis of many more samples. The CCL3L1 CNV has no simple relationships with flanking SNPs, but the availability of CN measurements for 5000 Biobank samples may allow the exploration of SNP-CN imputation, to ask whether phased haplotypes could form the basis of partial SNP-CNV imputation, or whether there is essentially complete linkage equilibrium between CNV and flanking SNPs. We have had some (unpublished) successes with the alpha-defensins using MOCShasper (Kato et al. 2008, Bioinformatics), and it could be useful to ask whether samples typed only at SNPs could be included in association analyses. Having said that, our own experience has been that the UK Biobank Axiom chip has sparse representation of SNPs around the amylase CNV, and it may be that the density of local SNPs near CCL3L1 may also be low because of the difficulties caused by local paralogy and CNV.

Minor typos (it would be much easier to specify these if the manuscript had line numbers):
Introduction, paragraph 3 has “In humans, Mip1alpha...”
There is variation throughout in the name given to the 32bp deletion in CCR5, and in particular whether the CCR5 component is italicised, and whether d or del is used.

In the Methods, section “CCL3L1 copy number estimation …”, I wasn’t sure what the intended meaning was for the sentence beginning “A prior information…”. Unless “an information” is a technical term I don’t know, presumably this should either be “Prior information…” or “A priori information…”. The following sentence should read “was fed into a Bayesian model…”.

References


Is the work clearly and accurately presented and does it cite the current literature?
Yes

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

**Competing Interests:** No competing interests were disclosed.

I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.