Modelling structural determinants of ventilation heterogeneity: a perturbative approach.

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Abstract

We have developed a computational model of gas mixing and ventilation in the human lung represented as a bifurcating network. We have simulated multiple-breath washout (MBW), a clinical test for measuring ventilation heterogeneity in patients with obstructive lung conditions. By applying airway constrictions inter-regionally, we have predicted the response of MBW indices to obstructions and found that they detect a narrow range of severe constrictions that reduce airway radius to between 10% -30% of healthy values. These results help to explain the success of the MBW test to distinguish obstructive lung conditions from healthy controls. Further, we have used a perturbative approach to account for intra-regional airway heterogeneity that avoids modelling each airway individually. We have found, for random airway heterogeneity, that the variance in MBW indices is greater when large-magnitude constrictions are already present, and that the indices become more sensitive to structural heterogeneity when already elevated. This method is a computationally efficient way to probe the lung’s sensitivity to structural changes, and to quantify uncertainty in predictions due to random variations in lung mechanical and structural properties.

Author summary

The multiple-breath washout (MBW) test is a clinical test of lung function that measures the efficiency of gas transport and mixing within the lung, and which has proven very sensitive in detecting early disease in cystic fibrosis (CF). In this paper we have developed a computational model of lung function to simulate air movement and gas transport in the lungs and generate MBW outcomes. We have used this to show why MBW is so sensitive to airway blockages similar to those encountered in CF.

Importantly, the model incorporates a new and computationally-efficient method that also allows us to account for uncertainty and randomness in lung structure and mechanical properties. This has been used to show how variability of MBW outcomes increases in disease states. The model provides a framework for modelling clinical data, where accounting for uncertainties in inputs is crucial in making clinically meaningful predictions.

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Introduction

The relationship between structure and function in the human lung is an important research area in physiology and medicine. Weibel’s casts of human lungs demonstrated that the bronchial tree can be treated as consisting of two functional regions \[1\]. First, the relatively rigid conducting part of the tree consists of dichotomously branching tubes that, on average, approximately follow the self-similar branching pattern given by Murray’s Law \[2\] allowing efficient transport of gas to the distal part of the lung. Second, the respiratory part of the lung (the acini) continues the dichotomously branching pattern but the tube lengths and diameters deviate from Murray’s law and airways terminate in alveolar sacs, where gas transfer with blood takes place \[3\]. The result is that flow velocity drops off very quickly in the acini and diffusion transports gas through the alveoli.

Ventilation heterogeneity (VH) arises in various obstructive lung diseases, such as cystic fibrosis (CF) and asthma, where inhaled gas is unevenly distributed in the lung, leading to poorer gas mixing efficiency \[4, 5\]. The severity of these conditions, in particular CF, is often quantified clinically using Multiple Breath Washout (MBW) tests \[6\]. MBW uses an inert tracer gas to quantify how effectively fresh air is turned over in the lung by measuring the tracer gas concentration and flow rate at the mouth. These data are used to compute clinically tested indices such as the lung clearance index (LCI) and phase-III slopes \[7\], which are indicators of VH.

Modelling the results of MBW tests accurately is challenging. First, modelling gas flows in a heterogeneous lung structure is a computationally expensive task. According to cast estimates there are on the order of $10^4 \rightarrow 10^5$ branches in the conducting zone \[8, 9\], and on the order of $10^7$ including the acinar ducts. There have been numerous approaches to resolve this problem, such as using a symmetric description with an effective diffusion coefficient to account for heterogeneous ventilation \[10\], compartmental models with asynchronous or asymmetric ventilation \[11, 12, 13\], modelling a single heterogeneous acinus \[14, 15\], and replacing the acini with well mixed units or symmetric models reducing the computation to $\sim 10^4$ branches \[16, 17, 18, 19, 20\]. Models have considered both heterogeneity in healthy lungs (such that average diameters and lengths converge to the healthy lung data) and pathological heterogeneity (where branches are constricted on average). Secondly, the majority of model parameters are difficult to measure experimentally, and are variable between subjects, which increases the uncertainty in predictions.

In this paper, we outline a perturbative approach to account for weak heterogeneity and the resulting uncertainty in ventilation within the lung. In this limit, gas transport can be estimated by a linear superposition of the solutions on trees with individual perturbations, greatly reducing computational expense. We simulate only the linearised response to perturbations and use these post-hoc to compute the response to different distributions of heterogeneity. The underlying physiological model we present assumes a linear elastic response of the alveoli, and Poiseuille flow throughout. However, the approach we present is generic and will be used in the future to quantify uncertainty and model the effects of heterogeneity in more complex constitutive models.

The perturbative approach can be used to describe lung structures with heterogeneity that is either deterministic (where the structural and mechanical properties are prescribed) or probabilistic (with properties described by multivariate distributions) within the same simulations. The probabilistic descriptions of lung structure means that we are able to directly estimate the variance of model outputs due to the parameter distributions and thus quantify uncertainty. We have applied this method to the particular case of MBW simulation.

In summary, the aim of this research was to model the complex mechanisms relating heterogeneity
and randomness in lung structure to MBW outcomes and their variability. To achieve this we had to develop an efficient method of accounting for weak structural heterogeneity that does not require simulating every airway and duct in the multi-scale lung structure individually.

**Methods**

**Computational Lung Model**

We have modelled the effects of inter-regional heterogeneity in the lung using a coupled network model of ventilation and gas transport in the airways and acinar ducts. To reduce model complexity, we have initially assumed that the airway tree can be approximated as completely symmetric within seven lung regions (see Figure 1); we refer to this as the mean-path model. Each region corresponds to a lobe or lobar compartment resembling the regions in Horsfield and Cumming’s lung model A [9]. We define the Strahler order of an airway by the maximum number of generations distal to it including acinar duct generations (counting from zero). Details of the model and parameters used can be found in S1 File and are summarised in S1 Table.

Gas flow on the airway network is calculated using a coupled set of ordinary differential equations (ODEs) that account for airway resistance (assuming Poiseuille flow), linear viscoelasticity of the acinar units and a uniform applied pleural pressure (similar to [17, 21, 18]). These ODEs are solved directly using the Eigen [22] factorisation routine ‘PartialPivLU’ in C++, as outlined in detail in S1 File §1.2. The concentration of inert gas on the network is then calculated using a one-dimensional advection-diffusion equation (S1 File §1.3) that accounts for transport into the alveolar sacs (similar to [23, 14, 15]). Transport in the acinar airways is included in this description as the details of its structure are important to accurately model gas transport [24, 25, 26, 27]. Within each lobar region the transport equation is identical to the ‘trumpet’ representation introduced in [28] (S1 File §1.4), with effective diffusion in the conducting region given by Taylor-like dispersion described in [29]. In the acinar region, pure molecular diffusion is modelled but with a modified airway cross-section, as described in [15]. The transport partial differential equations (PDEs) are discretised using a finite volume method (detailed in S1 File §2), and solved iteratively using the Eigen [22] ‘BiCGSTAB’ routine.

This lung model was used to simulate a multiple-breath washout (MBW) test by modelling the transport of an inert gas concentration $c$ on the lung network, where $c$ was assumed to be initially uniformly distributed. We selected tidal volume, functional residual capacity (FRC), airway deadspace volumes and lung elasticity values that are representative of a healthy adult male (see S1 Table). The MBW test measures FRC from the total inert gas exhaled over the test [7], which we label as $V_{\text{approx FRC}}$. The lung clearance index (LCI) value is taken to be the interpolated number of lung turnovers (exhaled volume in units of $V_{\text{approx FRC}}$) required to reduce the concentration (measured at the mouth at end of exhalation) to 2.5% of its initial value. LCI is a widely tested clinical measure of ventilation heterogeneity [30, 31], with healthy values generally in the range 6–8 and larger values indicating increased heterogeneity. Interpolation is necessary to measure small changes in LCI that are below the resolution of clinical LCI measurements, which are restricted to the set of end-tidal volumes (see S1 File §3). Phase-III indices measure ventilation heterogeneity through the slope of the measured gas concentration versus exhaled volume during individual breaths, and their interpretation is informed by numerical modelling [32, 33, 34]. In this paper we focus on $S_{\text{cond}}$, the linear gradient of the normalised phase III slopes, measured according to the clinical guidelines for MBW in [7].
more recent development in the field comes from hyperpolarised helium MRI imaging of the inert gas concentration in lungs [35]. Using this technique, fractional ventilation of lung acini is also quantified by calculating the dilution rate of the inert gas concentration between successive breaths at end-tidal volume [36]. Heterogeneity in fractional ventilation correlates strongly with increased LCI, but the MRI images can also identify changes not picked up by MBW indices, as well as provide important information about the spatial distribution of ventilation [6]. While the network model presented here cannot predict the spatial structure of FV distributions measured in MRI, it can predict distribution within each model region.

**Perturbative method and application to structural heterogeneity**

Building on this model, we have used perturbation theory to calculate the changes in gas concentration due to small variations of the properties of a single airway or acinus in a mean-path. This
involves calculating the linearised response of the simulation variables with respect to the perturbation magnitude, modelled by the dimensionless parameter $\epsilon \ll 1$. For example, if a given airway has cross-sectional area $a$ in the baseline model, it becomes $a(1 + \epsilon)$ in the perturbed case, and the response of all other simulation variables, such as gas concentration $c$, are computed to linear order in $\epsilon$. For each perturbation considered, the number of separate paths that need to be simulated grows linearly with the airway generation number.

In “Intra-regional heterogeneity”, we have used this method to model intra-regional heterogeneity in airway sizes, by assuming they are randomly distributed about their mean-path values, and that the variance of this distribution is small enough that we can apply perturbation theory as detailed in S1 File §1.5. In the simple case where the perturbations are modelled as variables in a multivariate Gaussian, the variance in model outputs can be computed as a sum of the covariance of inputs weighted by their respective linear sensitivities (see S1 File §3.) This gives a computationally efficient method to relate weak intra-regional heterogeneity to variance of model outputs that relies only on each constricted case being simulated once.

We have modelled uncorrelated, normally distributed perturbations to the area and length of the conducting airways. We have assumed that the coefficient of variation of these parameters, $\sigma_a$ and $\sigma_l$, is independent of generation and path, (see S1 File §3.1 for a mathematical description of this distribution). We have also introduced structurally correlated uncertainty in geometry within each Independent Structurally Correlated independent case being simulated once.

![Figure 2](image_url)

Figure 2: Sketch of randomly distributed airway sizes, shading (from dark to light) indicates airway size relative to generational average. In the independent case, variations in branch size are uncorrelated. In the correlated case larger than average airways are likely to beget airways that are also larger than average, resulting in an increased uncertainty in the size of the most distal airways. For simplicity, all airways in this sketch have fixed aspect ratio (the case $\rho_{al} = 1$). When $\rho_{al} = 0$, airway length and area are independent.

mean path. The assumption made here is that the deviation of each branch from the mean-path value is normally distributed around that of its parent branch (see Figure 2), again with coefficients of variation $\sigma_a$, $\sigma_l$. The area and length deviation of a branch is itself taken to be correlated with coefficient $-1 \leq \rho_{al} \leq 1$. In what follows we present the case $\sigma_a = 2\sigma_l$ (in the linear limit radius and length have the same coefficient of variance), for $\rho_{al} = 0$ (no area and length correlation) and $\rho_{al} = 1$ (fixed aspect ratio). A complete description of this case is given in S1 File §3.2. This case is similar to the random structure generated used for particle deposition calculations in [37], whereby airway sizes are generated according to the size of the parent airway.

Finally, we have used these results to compute the probability density of any acinus in the model.
having a given FV value (averaged over the whole MBW test). In the limit of a large number of acini, the
distribution of FV values in any given lung model realisation will tend towards this distribution
(see S1 File §3 for further details).

Model validation

Validation studies to test the accuracy and precision of the numerical simulations can be found accom-
panying the source code at [45]. In these we have tested that the code is suitably converged for the
choice of model time-step and space-step, and that inert gas volume is conserved. We also tested that
the perturbative model converges exactly to the mean-path model in the limit of small perturbations.

Results

The healthy baseline model (Figure 1) assumed that all regions have the same airway sizes and
mechanical properties. This resulted in a homogeneous distribution of gas, with the only asymmetry
originating from the arrangement of the proximal airways supplying the lobar regions, and the number
of generations within each region. In this case the simulations showed little regional difference in
fractional ventilation and each lobe contributed proportionately to the washout. The baseline values
of the MBW indices (using SF$_6$ as the tracer gas) were $\text{LCI} = 5.16$, $\text{S}_{\text{cond}} = 1.92 \times 10^{-4}$L$^{-1}$ and
$\sqrt{\text{FRC}}(\text{approx}) = 3.00$L (to 3 s.f.). Using N$_2$ the LCI reduced slightly to 5.04, due to better gas mixing and
a more proximal diffusion front. S1 Figure compares the linear sensitivity of LCI to airway size for
both gases (computed using the perturbative method), demonstrating that the key differences between
the two gases are in the acinar generations, where the diffusion front is located.

Inter-regional heterogeneity

Figure 3 (solid lines) shows the effect of constricting airways in the right-middle (RM) lobe at three dif-
ferent depths. Each case simulates localised bronchoconstriction, with all airways in a given generation
range (proximal, central or distal) reduced in cross-section by the same fraction, approximating the
pathophysiology of asthma or CF. A marked response in all three MBW indices was evident for radius
constrictions above circa 70%. The responses of LCI and $\text{S}_{\text{cond}}$ to airway constrictions were strongly
correlated, peaking at approximately 80% constriction of the radius before dropping back to baseline
values (Figures 3(a) and (b)), similar to the results reported in [16, 20]. This response was dominated
by the resistance of the constricted airways (see S2 Figure) which resulted in under-ventilation of
the affected regions, as well as asynchrony in emptying (see S1 Video and S2 Video). This response
was effectively independent of constriction depth and dropped off at larger constrictions, where these
regions became essentially unventilated and thus undetectable at the mouth. This is shown in Figure
3(e) by the reduction in measured FRC volume of approximately 10% (295 ml) of the lung volume.

It is a simplification to assume that constrictions or blockages would be localised to a single lobe.
However, randomly distributed constrictions applied to families of airways at each depth that feed the
same fraction (10%) of the lung volume, resulted in a very similar response due to the homogeneity
of the baseline case (see Figures 3(b), (d) and (f)). The response was weaker than the localised case
for a more distal heterogeneous distribution of constrictions, and dropped off more gradually with
very large constriction sizes (>80% radius constriction). In this case, 10% of the airways from the
most proximal generation (Strahler orders 19,15,11 for proximal, central and distal cases respectively)
were selected uniformly randomly, and constrictions were applied to these and the three generations
Figure 3: Relationship between constriction strength (% radius reduction) and MBW indices (a)-(b) LCI, (c)-(d) $S_{\text{cond}}$ and (e)-(f) measured FRC volume $V_{\text{FRC}}^{\text{approx}}$ applied to airways feeding 10% of the lung acinar volume. Three different depths were tested, corresponding to Strahler order 19-16 (proximal, black squares), 15-12 (central, red circles), and 11-8 (distal, blue diamonds), where all branches constricted were taken to be directly descended from the most proximal in all cases. (a), (c) and (e) show simulations where constrictions were all localised within the right-middle lobe, whereas (b), (d), and (f) show realisations where the positions of the constrictions were uniformly randomly distributed throughout the lung. Example animations of localised and random constrictions are shown in S1 Video and S2 Video respectively.

of airways descended from them. Once constrictions had been applied to the full network, symmetric sub-trees were re-identified and replaced with mean-paths (following §1.4), as can be observed in the network structure in S2 Video. Note that these results were computed through direct simulation, and not using the perturbative approach.

To conclude, we have found that MBW indices detect a restricted range of severe airway constrictions, which our results predict to be most sensitive when airways are between 10% - 30% of their original radius.
Intra-regional heterogeneity

Figure 4 shows the predicted uncertainty in MBW indices due to airway heterogeneity. Two types of airway heterogeneity are presented: first where changes to airway lengths and areas are generated completely independently; and second where the areas and lengths are correlated with changes in the parent airways (see Figure 2 and S1 File §3 for further details). In general, the standard deviations of LCI and $S_{\text{cond}}$ increased when airway sizes were correlated with their parent branch. Furthermore, the standard deviations showed a similar response to constrictions as the indices themselves, meaning that there was increased uncertainty due to airway heterogeneity when the indices were elevated. There was a small drop in LCI variance at $\sim$ 80% radius constrictions, corresponding to the stationary point of the LCI curve in Figure 3(a)-(b). This means that the uncertainty showed similar behaviour to the magnitude of the gradient of the curve in Figure 3(a)-(b).

![Figure 4](image)

Figure 4: Standard deviation in (a)-(b) LCI and (c)-(d) $S_{\text{cond}}$ vs. constriction strength (for constrictions confined to RM lobe). Results were computed for independent normally distributed perturbations (black squares) and structurally correlated perturbations (see S1 File §4.2) with $\rho_{al} = 0$ (diameter and length uncorrelated, red circles) and $\rho_{al} = 1$ (fixed diameter-length ratio, blue diamonds). (a) and (c) show the results for Strahler orders 19-16 (proximal) and (b) and (d) for Strahler orders 11-8 (distal, dotted line, diamonds). Constrictions were applied to all airways in the RM lobe within these generation ranges as in Figure 3. In all cases $\sigma_a = 0.2$ and $\sigma_l = 0.1$. Note the logarithmic scale on the vertical axes.

Figure 4 also shows different responses depending on constriction depth. Independently distributed airway heterogeneity has less effect on standard deviation of the indices when constrictions are more distal. However, when parent-daughter airway sizes were correlated, the effect on LCI and $S_{\text{cond}}$ standard deviations was similar regardless of constriction depth. This implies that this type of structured heterogeneity, whereby uncertainty in airway size grows with each generation, enhances the
contribution of the smaller airways to the overall uncertainty.

Figure 4 also compares the case where airways retain a fixed length-to-diameter ratio, but there is heterogeneity in overall airway size. In this case, the standard deviations of LCI and $S_{\text{cond}}$ did not increase by as much at severe constriction strengths. Area and length increases have opposite impacts on airway resistance, and therefore we observed opposite effects LCI and $S_{\text{cond}}$ at severe constrictions (see S2 Figure). Thus correlated area and length changes ($\rho_{al} = 1$) did not increase the uncertainty as much as uncorrelated ($\rho_{al} = 0$) changes when severe constrictions were present.

Finally, the perturbative simulations also showed that sensitivity of the MBW indices to changes in lung elastance was increased at large constriction strength (see S3 Figure). Also, even though the constrictions were localised to the RM lobe, this increased sensitivity was observed for changes in elastance in all of the lobes.

To conclude, when the MBW indices are elevated due to severe airway constrictions, they were more sensitive to heterogeneity in both airway size and other mechanical properties such as elastance. This indicates that values measured experimentally are likely to be more variable in the presence of severe bronchoconstriction or mucus plugging.

Global Heterogeneity: Fractional Ventilation Distribution

Figure 5 shows the distribution of acinar fractional ventilation (FV) values at different constriction strengths, and for different distributions of airway heterogeneity. The dotted vertical lines in Figure 5(b)-(d) show the FV values for each region in the baseline model (no linear perturbations), while the continuous distributions show the predicted FV values by the perturbative model. As the fractional ventilation in the RM lobe decreased (and LCI increased) the distribution of FV in this lobe also broadened (Figures 5(b) and (c)), before it narrowed again at very large constrictions (Figure 5(d)). This shows that the local FV was most sensitive to airway heterogeneity in the same constriction strength range as the MBW indices. When there was fixed airway length-to-diameter ratio) the FV distribution was narrower in the constricted lobe (RM), where airway resistance dominates, and broader in the other regions, where airway dead-space volume is the dominant factor. The results clearly demonstrate a link between the width of the FV distribution and the resulting model uncertainty in MBW indices due to airway heterogeneity.

A key finding is that the unconstricted mean-paths were relatively unaffected, and remained fairly insensitive to airway heterogeneity in those paths. Nonetheless, there is a small drop in FV in the right-lower lobe, which can be explained by a pendelluft effect where gas from the right-middle lobe is re-inspired into the right-lower lobe due to the asynchronous nature of the ventilation (see S1 Video).

In summary, the FV distributions show that severe airway constrictions result in a much broader distribution of gas turnover in the affected lung region (assuming some randomness in airway geometry). Aside from the noted pendelluft affects, the distribution of FV due to airway randomness remained largely unchanged, highlighting the parallel nature of the airway network structure.

Discussion

In this paper we have introduced a perturbative approach that accounts for the uncertainty in predictions due to random variation in the lung’s geometrical and mechanical properties. Our results showed that $S_{\text{cond}}$ and LCI responded in a highly correlated manner to bronchoconstriction, whether this was localised to a single lobe or distributed across the lung (Figure 3). The response was notably non-
Figure 5: Whole lung fractional ventilation distributions for (a) 0% (b) 71.5% (c) 81.5%, and (d) 91.5% constrictions to the radius of the airways in the central airways of the RM lobe (central, Strahler orders 15-12). Results for independent random perturbations (solid black lines) and structurally correlated random perturbations with $\rho_{al} = 0$ (dashed red lines) and $\rho_{al} = 1$ (fixed diameter-length ratio, dotted blue lines). For visibility, $\sigma_a = 0.5$ and $\sigma_l = 0.25$ was used for the independent case, whereas $\sigma_a = 0.1$ and $\sigma_l = 0.05$ was used for the correlated case. The vertical dotted lines indicate the mean FV values in each lobar region. In (b), (c) and (d) the regions are labelled as in Figure 1.

linear, showing a high sensitivity to large-magnitude constrictions (~80% reduction in radius) before returning to baseline values at even larger constrictions. The sharp response is due to the inverse-fourth-power dependence of airway resistance (approximated here by Poiseuille flow) to changes in airway radius. This suggests a mechanism to explain the success of MBW indices to differentiating obstructive lung conditions such as asthma and CF, where airway inflammation and blockage is a typical feature, from healthy volunteers where such narrow airways are unlikely to be present in large
numbers. We also found that LCI and $S_{\text{cond}}$ did not respond as strongly if constrictions were randomly distributed in the distal conducting airways, because in this case the well- and under-ventilated lung units branch from one another more distally (where advection is less dominant over diffusion) than the case where constrictions were localised.

This weak response to small airway geometry deviations motivated the use of perturbation theory to account for heterogeneity in the underlying airway structure. Using this, we found that the uncertainty in predictions due to weak heterogeneity in the structure was greatly amplified when LCI and $S_{\text{cond}}$ were elevated due to constrictions (where we considered constrictions confined to a single lobe, see Figure 4). When the perturbations were independently distributed, the variance was greater if there were few proximal constrictions (two in this case) rather than numerous distal constrictions ($2^9$). This is because the indices are most sensitive to geometrical changes in the perturbed airways, which are more likely to average out due to the large number of constrictions in the latter case. On the other hand, fluctuations from the mean are much more likely to occur when there is a small number of constrictions. This uncertainty is amplified further by including structural correlations that account for the inherited nature of airway sizes (e.g. larger-than-average airways on average branch into larger-than-average airways). We found that the smaller airways appeared to contribute equally to the variance in indices in this case, because more proximal fluctuations from the mean were propagated down the airway tree.

More broadly, these results suggest that elevated MBW indices induced by airway constrictions are more variable in general, which is observed experimentally through the increased variance in LCI measured in CF and asthma patients [46], as well as an increased sensitivity of LCI to posture which affects the distribution of ventilation due to gravity [47, 48]. This latter affect can be understood by perturbing acinar elastance, which affects their rate of ventilation. Figure 5 confirmed that the magnitude of the sensitivity of the MBW indices to elastance perturbations was much larger when airways were pathologically constricted (even when the perturbations were in different lung regions to the constrictions). The former effect is more complex, as inter-subject variability is likely to be larger due to variations in disease progression and expression, which would be expected to dominate over the effects of heterogeneity in airway sizes. Additionally, in the artificial case considered here, the constriction strengths are assumed uniform and hence extra uncertainty would be introduced by including heterogeneity in constriction strength too.

Finally, we used the perturbative approach to compute probability distributions of acinar fractional ventilation values (Figure 5). The FV distributions predicted are generally narrower than measured experimentally [36]. This is in part due to the simplified nature of the mean-path model and the assumption of weak heterogeneity. Furthermore, acini sizes are also variable which directly affects their FV values, while gravitational effects also play a role [17]. However, the response we observe is indicative of the effects of heterogeneity in airway structure alone, and demonstrate the relationship between variation in structure and the distribution of FV within a lung. We saw that, generally, the FV distribution was much broader in the lung region affected by the constrictions, which is consistent with the increased variance in MBW indices predicted (Figure 4). These calculations also showed that the unconstricted regions are relatively unaffected by the presence of the blockage, due to the parallel nature of the lung structure.

The underlying model is simple, considering only seven distinct lung regions and assuming that the airway and acinar structures are symmetrical in each. However, the model is sufficient to predict some generic relationships between lung structure and the MBW indices, in particular LCI and $S_{\text{cond}}$. 
The LCI values predicted by the model are low compared to those measured experimentally [10], but similar to those simulated in more detailed airway tree models [39, 20]. The phase-III slopes are practically zero in the absence of constrictions, whereas in a healthy lung airway asymmetry and acinar duct asymmetry both contribute to positive slopes [39]. These differences can in part be attributed to the idealised nature of this model, which assumes complete symmetry in the acinar structure. This means that mixing efficiency in the alveolar zone is better than is likely in reality, and LCI is therefore lower.

The perturbative method addresses this effect of intra-regional airway asymmetry, which is not present in the baseline model, but is valid only for small deviations of properties from the mean. As this method measures the linearised response of the model variables to changes in structure or mechanics, it misses non-linear behaviour, which can become dominant at increasing perturbation magnitude. Additionally, the number of trees one has to simulate to compute all of the linear sensitivities increases with the number of mean-paths in the baseline model. Thus there is a balance to be struck between the resolution of the baseline model (i.e. how many lung regions the mean-path model is split into) being sufficient to simulate realistic ventilation heterogeneity and computational efficiency. Nonetheless, the perturbative approach gives a unique insight into the relation between structural changes and lung function.

Other assumptions made in the model are likely to affect predicted MBW outcomes. Most significantly, we have neglected the effects of gravity/posture on inter-regional variation, as well as mechanical coupling of the lung units, which are both predicted to affect fractional ventilation and MBW indices in simulations [10, 17, 41, 42] and experiments [43, 44, 48]. The lack of mechanical coupling means that the predicted asynchrony between lung regions may be exaggerated compared to reality, which could indicate why the range of predicted $S_{\text{cond}}$ values is notably wider than the increase measured between healthy volunteers and CF patients [50]. Furthermore, air flow has been modelled by the Poiseuille relation in all airways, meaning that the effects of inertia [38] and turbulent flow are neglected. Finally, we have not include the effects of gas exchange on inert gas transport, as it is thought to be negligible (except in the case where nitrogen is used as the MBW tracer gas [49]). The limitations imposed by these assumptions are discussed in more detail in S1 File.

**Conclusion**

The strength of the perturbative approach introduced in this paper is two-fold. First, it reduces simulation complexity, as the maximum number of terminal airways to be simulated scales linearly with generation rather than exponentially. Second, once the linear sensitivities are calculated for a given simulation, they can be combined in numerous ways to investigate the effects of different patterns of heterogeneity. This avoids the need to randomly generate many realisations of a higher-resolution model, as would be required if using Monte-Carlo sampling.

In future, this approach will be further developed to quantify uncertainty in more realistic lung models that are directly informed by imaging data. The principles outlined here will enable a systematic approach that quantifies uncertainty due to both the intrinsic complexity of lung structure and the additional effects of obstructive lung disease or gravity.
Acknowledgments

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Supporting information

S1 Table  List of parameters and values used in simulations.

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<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
<th>Values used</th>
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<tr>
<td>$V_D$</td>
<td>Total volume of conducting airways.</td>
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<td>$V_{mouth}$</td>
<td>Total volume of mouth cavity.</td>
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<td>Airway length scaling in mean-path conducting branches.</td>
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<td>$\lambda_{acin}$</td>
<td>Duct length scaling in acinar branches.</td>
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<td>Number of acinar airway generations.</td>
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</tr>
<tr>
<td>$\Phi_j$</td>
<td>Density of alveolar sacs. in acinar generations.</td>
<td>0 if $j &lt; N_{z}^{cond}$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.2 if $j = N_{z}^{cond}$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.4 if $j = N_{z}^{cond} + 1$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.7 if $j = N_{z}^{cond} + 2$</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1 if $j &gt; N_{z}^{cond} + 2$</td>
</tr>
<tr>
<td>$\phi$</td>
<td>Phenomenological parameter, see equation (10) in S1 File</td>
<td>0.5 [15]</td>
</tr>
<tr>
<td>$K$</td>
<td>Elasticity of the whole lung.</td>
<td>5 cm H$_2$O L$^{-1}$ [52, 53]</td>
</tr>
<tr>
<td>$R_{acin}$</td>
<td>Cumulative resistance of acini.</td>
<td>0.2 cm H$_2$O s L$^{-1}$ [54, 52]</td>
</tr>
<tr>
<td>$R_{UA}$</td>
<td>Resistance of the upper airway.</td>
<td>0.6 cm H$_2$O s L$^{-1}$ [54, 52]</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Air viscosity at 37$^\circ$C.</td>
<td>1.93 $\times 10^{-7}$ cm H$_2$O s</td>
</tr>
<tr>
<td>${d, l}$</td>
<td>Trachea, Right Main Bronchus, Left Main Bronchus, Right Intermediate Bronchus, Right Lower Lobar Bronchus, Left Lower Lobar Bronchus</td>
<td>{1.6, 10}</td>
</tr>
<tr>
<td></td>
<td></td>
<td>{1.11, 2.2}</td>
</tr>
<tr>
<td></td>
<td></td>
<td>{1.2, 5}</td>
</tr>
<tr>
<td></td>
<td></td>
<td>{0.89, 2.6}</td>
</tr>
<tr>
<td></td>
<td></td>
<td>{0.64, 0.8}</td>
</tr>
<tr>
<td></td>
<td></td>
<td>{0.8, 1.1}</td>
</tr>
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</table>
S1 Figure  Linear fractional change in LCI due to a single perturbation in area ((a) and (c)) and length ((b) and (d)) scaled by number of branches in that generation $N_b$. The airway generation is plotted in terms of its Strahler order (i.e. its generation counting up from zero at the bottom of the tree). The vertical dashed line indicate the terminal bronchiole separating the acinar (Strahler orders 0-8) and conducting (>9) generations. (a)-(b) Healthy lung model using SF$_6$ (molecular diffusivity 0.105cm$^2$ s$^{-1}$). (c)-(d) Healthy lung model using N$_2$ (molecular diffusivity 0.225cm$^2$ s$^{-1}$). Coloured symbols distinguish perturbations in the seven lobar regions. The LCI sensitivities in the conducting region (right of the vertical dashed line) are approximately identical for area and length perturbations in both cases, as this is a response to the increase in dead-space volume. For SF$_6$ the sensitivities in the acinar region (left of the vertical dashed line) are inverted for length and area perturbations, most notably around the diffusion front (approximately Strahler order 4). Thus LCI is sensitive to geometry changes that affect diffusion in the acinus when using the less diffusive SF$_6$, but not N$_2$. 
S2 Figure  Linear sensitivities to geometry perturbations in a model realisation with (a)-(b) 74% and (c)-(d) 81.5% constrictions in radius to the RM lobe. Scaled sensitivities (as in S1 Figure) w.r.t. area (filled markers) and length (open markers) of the airways are shown for the RM lobe only for fractional LCI change and absolute change in $S_{\text{cond}}$. Results were plotted for different depths of constriction: proximal (Strahler orders 16-19, black squares), central (Strahler orders 12-15, red circles) and distal (Strahler orders 8-11, blue diamonds). The sensitivities are scaled by the number of airways in the corresponding Strahler order of the RM lobe. The scaled sensitivities are much larger in the constricted airways, as the response is most sensitive to their resistance (note the difference in scale to S1 Figure). Since airway resistance scales as length/area$^2$, the area sensitivities are approximately a factor $-2$ of the length sensitivities. The sign of the sensitivities changes between the two constriction strengths because they lie either side of the maximum values of LCI and $S_{\text{cond}}$ in Figure 3(a) and (b).
S3 Figure  Sensitivity of LCI to changes in elastance of an acinus in each mean-path vs. constriction strength of the central airways in the RM lobe. Linear sensitivities were scaled by the number of acini \( N_b \) in the perturbed lobar region. The sensitivity increased in size by an order of magnitude upon increasing constriction size. As constriction strength was increased, an increase in elasticity of the RM lobe decreased flow to that lobe, further increasing ventilation heterogeneity and LCI. Meanwhile increases to elasticity in other lobes increased the flow to RM and decreased LCI. Once the constriction strength is > 75% the effect reversed because increases in the flow to the RM lobe released trapped gas that would have otherwise been undetectable by MBW.

S1 Video  Inert gas concentration on lung network for various constriction strengths to the central airways of the RM lobe (% reduction in radius as shown). Vertical direction is the distance from the mouth, while horizontal distances have no physical meaning and are set for visibility. Time scale 1:4 (each second of video corresponds to 4 seconds of washout).

S2 Video  Inert gas concentration on lung network at varying constriction strengths in the central airways distributed throughout the lung (% reduction in radius as shown). Vertical direction is the distance from the mouth, while horizontal distances have no physical meaning and are set for visibility. Time scale 1:4 (each second of video corresponds to 4 seconds of washout).

S1 File  Supplementary text containing further details of the methodology.

References


[38] Pedley TJ, Schroter RC, Sudlow MF. The prediction of pressure drop and variation of resistance within the human bronchial airways. Respir Physiol. 1970;9(3):387–405. doi:10.1016/0034-5687(70)90094-0


