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- 2 parameters derived from hyperpolarized gas lung ventilation MRI in healthy
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- edited and revised manuscript, approved final version of manuscript
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- 17 version of manuscript
- Bilal A. Tahir: Analyzed data, edited and revised manuscript, approved final version
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Running head: Multiple lung inflation level imaging using HP gas MRI

ABSTRACT: In this study, the effect of lung volume on quantitative measures of lung ventilation was investigated using MRI with hyperpolarized ³He and ¹²⁹Xe. Six volunteers were imaged with hyperpolarized ³He at five different lung volumes (residual volume (RV), RV+1L, functional residual capacity (FRC), FRC+1L and total lung capacity (TLC)), and three were also imaged with hyperpolarized ¹²⁹Xe. Imaging at each of the lung volumes was repeated twice on the same day with corresponding ¹H lung anatomical images. Percentage lung ventilated volume (%VV) and variation of signal intensity (heterogeneity score, H_{score}) were evaluated. Increased ventilation heterogeneity, quantified by reduced %VV and increased H_{score}, was observed at lower lung volumes with the least ventilation heterogeneity observed at TLC. For ³He MRI data, the coefficient of variation of %VV was less than 1.5% and less than 5.5% for H_{score} at all lung volumes, whilst for $^{129}\mathrm{Xe}$ data the values were 4% and 10% respectively. Generally, %VV generated from ¹²⁹Xe images was lower than that seen from ³He images. The good repeatability of ³He %VV found here supports prior publications showing that percentage lung ventilated volume is a robust method for assessing global lung ventilation. The greater ventilation heterogeneity observed at lower lung volumes indicates that there may be partial airway closure in healthy lungs and that lung volume should be carefully considered for reliable longitudinal measurements of %VV and H_{score}. The results suggest that imaging patients at different lung volumes may help to elucidate obstructive disease pathophysiology and progression.

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KEYWORDS: Imaging; MRI; Hyperpolarized gas; Lungs; Inflation;

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63	NEW AND NOTEWORTHY: We present repeatability data of quantitative metrics
64	of lung function derived from hyperpolarized helium-3, xenon-129 and proton
65	anatomical images acquired at five lung volumes in volunteers. Increased regional
66	ventilation heterogeneity at lower lung inflation levels was observed in the lungs of
67	healthy volunteers.
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69	ABSTRACT WORD COUNT: 246/250
70	NEW AND NOTEWORTHY WORD COUNT: 43/75
71	MANUSCRIPT WORD COUNT:

FIGURE COUNT: 9

TABLE COUNT: 7

Introduction

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Hyperpolarized (HP) gas ventilation-weighed magnetic resonance imaging (MRI) 76 77 allows the visualization of gas distribution within the lung and has been shown to 78 detect early lung disease in patients with cystic fibrosis and normal spirometry (4, 19) 79 and in the lungs of smokers (33). Additionally, it has been used to assess the response 80 to treatment in patients with asthma (13, 29), to longitudinally assess patients with 81 chronic obstructive pulmonary disease (17) and has been shown to be clinically 82 feasible for assessing lung function in children (1). 83 HP gas and proton anatomical (¹H) lung magnetic resonance imaging (MRI) can be 84 85 combined to quantify lung ventilation using percentage lung ventilated volume 86 (%VV) or its counterpart the ventilation defect percentage (%VDP) (33), both of 87 which have been widely adopted as simple and robust image-derived metrics. %VV is 88 the ratio of the ventilated lung, defined from HP gas ventilation-weighted images to the thoracic cavity volume, defined from the ¹H anatomical images (16, 33). Previous 89 90 work has shown improved repeatability of %VV when the anatomical image is 91 acquired in the same breath-hold as the HP gas ventilation-weighted image (12). 92 93 Ventilation heterogeneity may be assessed by using the H_{score} metric developed by 94 Tzeng et al. (31), which calculates the variation of signal intensity in a kernel around 95 a given voxel as the standard deviation divided by the mean (e.g. Figure 1) (23, 31). 96 97 The clinical standard for assessing lung volumes is body plethysmography (22, 32), 98 whilst changes in forced expiratory volume in one second (FEV₁) and forced vital 99 capacity, measured using spirometry, are used as clinical markers for lung function

decline in certain diseases (22, 32). However, patient coaching of inhalation from a bag of gas rather than spirometric gating is generally used to achieve the lung volumes for HP gas MR imaging, which may lead to variability in lung volumes as will the ability of the patient to inhale the entire contents of the bag of gas being used. The most frequently used lung volume is functional residual capacity plus 1 liter (FRC+1L) (6-8, 16, 18, 27, 34). However, if a 1L bag is inhaled from FRC in smaller patients this volume may be close to total lung capacity (TLC) and thus understanding the effect of lung inflation level on these image-derived metrics is important.

Previous work by Muradyan et al. (23) analyzed the effect of inhalation of HP xenon-129 (129 Xe) from residual volume (RV) in healthy volunteers and sub-RV in elite divers by acquiring coronal projection images with an in-plane resolution of 4.7mm x 9.4mm. Muradyan et al. calculated the global H_{score} in the ventilated regions of the image, and found that when the elite divers inhaled low volumes of gas (0.9L and 0.4L respectively) compared to larger volumes of gas (1.3L and 0.9L respectively) from sub-RV, increased heterogeneity was seen in the images, consistent with punctate reopening of some airways that were closed at sub-RV. Marshall et al. (20) carried out preliminary work demonstrating the effect of airway opening between FRC+1L and TLC using HP ³He imaging showing decreased heterogeneity and increased %VV at TLC when compared to FRC+1L. With these studies demonstrating important mechanisms at work in healthy controls and patients it is clear that understanding the effect of lung inflation on quantitative metrics derived from HP gas and ¹H anatomical MRI is an important step in moving these techniques forward into standard clinical practice.

125	Historically, noble gas MRI studies have made use of HP helium-3 (³ He); however,
126	with the rising cost and scarcity of ³ He, the focus of the pulmonary imaging
127	community is switching to the use of HP ¹²⁹ Xe (18, 27) where differences in metrics
128	have been reported due to the differences in diffusivity and the achievable signal of
129	¹²⁹ Xe MRI. Thus the aims of this study were to use both HP ³ He and ¹²⁹ Xe MRI to:
130	1. assess the effect of different lung inflation levels on the HP gas image derived
131	metrics %VV and H _{score} .
132	2. assess the repeatability of %VV and H_{score} from two same-day imaging
133	sessions.
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135	Materials and methods
136	<u>Subjects</u>
137	The study was performed with national research ethics committee approval and with
138	informed consent from all volunteers. Six volunteers (all male) were recruited for this
139	study with the only criterion being that subjects were suitable for MRI and had no
140	known respiratory complications. Two volunteers were former smokers, two were
141	occasional smokers and two were never smokers. Table 1 shows the subject
142	demographics.
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144	Study protocol
145	Spirometry was performed to international standards (32) to ensure subjects had were
146	defined as spirometrically free from respiratory conditions.

All ³He imaging was carried out on a GE HDx 1.5T MRI scanner (GE Healthcare, Milwaukee, WI, USA) using a ³He transmit-receive flexible chest coil (Clinical MR Solutions, Brookfield, WI, USA). ³He was polarized using a commercial polarizer (GE Healthcare, Amersham, UK). HP ³He 3D balanced steady state free precession and ¹H spoiled gradient echo images were acquired in the same breath (12) at five different lung volumes: RV, RV+1L, FRC, FRC+1L and TLC. For ¹²⁹Xe imaging, the gas was polarized using a home-built polarizer (24) and images were acquired using a ¹²⁹Xe transmit-receive flexible vest coil (Clinical MR Solutions, Brookfield, WI, USA) and the ¹H system body coil at five different lung volumes, as with ³He imaging. ¹²⁹Xe and ¹H images were acquired in separate breath-holds as previously described (27, 28) and this was due to the longer acquisition time of the ¹²⁹Xe scan. Note that only a subset of the volunteers (V2, V3 and V6) were scanned using HP ¹²⁹Xe and separate-breath ¹H imaging as a feasibility study as some participants were no longer available to be scanned. A 1L mixture of hyperpolarized gas and nitrogen was used as it is the most commonly used volume in adults (6-8, 16, 18, 27, 34).

For the breathing maneuvers (Figure 2), volunteers were coached and instructed to breathe within the scanner by a pulmonary physiologist. During imaging, breathing maneuvers started with inhalation of the contents of the 1L bag from FRC, except for imaging at RV+1L where volunteers first exhaled to RV. To acquire images at TLC, volunteers inhaled room air to maximum lung capacity after the inhalation of 1L of gas from the bag. For imaging at FRC, volunteers inhaled the contents of the 1L bag from FRC and then exhaled back to FRC. For RV imaging, volunteers inhaled the contents of the 1L bag from FRC and then exhaled to RV. Gas doses were increased for the exhalation maneuvers and for imaging at TLC with the aim of ensuring

sufficient signal for imaging, and prior to exhalation participants held their breath for 5 seconds to allow the gas to diffuse into the peripheral lung. Inhaled gas doses are given in Table 2; note that images were also acquired in the order presented in Table 2.

For ³He acquisitions, subjects were scanned twice on the same day, with a 10 to 20-minute break (remaining supine within the scanner) in between imaging sessions. ³He imaging sessions lasted 20-30 minutes on average. For ¹²⁹Xe imaging, subjects were scanned twice on the same day, with a 20 to 40-minute break between imaging sessions and were removed from the scanner during this break. ¹²⁹Xe imaging sessions lasted 35-45 minutes on average, due to limitations imposed by gas polarization time.

Image analysis

Thoracic cavity volume (TCV) and ventilated volume (VV) were extracted from the 1 H anatomical and HP gas ventilation images, respectively, using the semi-automated segmentation method based on spatial Fuzzy C-means thresholding previously described (15). Percentage lung ventilated volume was calculated according to $\%VV = (VV/TCV) \times 100$.

Ventilation heterogeneity was assessed using a modified version of the H_{score} method previously described (31). Images were subsampled from 256x256 voxels in-plane to 128x128 voxels, resulting in an apparent image resolution of ~3.2x3.2x5mm for 3 He images or ~3.2x3.2x10mm for 129 Xe images. To avoid partial volume effects at the edge of the ventilation-weighted images, the TCV mask was eroded by 1 pixel, and

the ventilation-weighted image was then multiplied by the VV mask and eroded TCV masks, with voxels outside of the VV and TCV masks being excluded from the local heterogeneity calculation. To generate maps of ventilation heterogeneity, a 3x3 voxel kernel (~9x9mm) was then passed over the images, centered on every voxel in the ventilated volume, to calculate the local variation of signal intensity (H_{i,j,k} at voxel i,j,k). H_{score} in this work was then defined as the median of the non-zero values of the local heterogeneity map rather than the mean as previously reported, as the histograms of H_{score} were not normally distributed. For images acquired at TLC, where there was clear signal dropout due to coil sensitivity coverage, VV and TCV masks were matched, i.e. where signal dropout occurred emulating a defect it was manually excluded on both the TCV and VV masks, in order to ensure that this did not cause increased H_{score} and decreased %VV.

Additionally, the mean H_{score} of the most posterior slice was compared to the mean H_{score} of the remaining image slices for each volunteer at each inflation level for the data acquired with ${}^{3}He$. The mean values were grouped by volunteer and lung volume and significant differences were assessed using either a paired t-test or Wilcoxon matched-pairs signed rank test depending on the normality of the data. This analysis was not carried out for ${}^{129}Xe$ data due to the reduced number of subjects.

Repeatability and statistical analysis

To assess the repeatability of %VV and H_{score} between session 1 (S1) and session 2

(S2), the coefficient of variation (CoV), Bland-Altman analysis (2), paired t-tests and

the repeatability limit were used. For CoV analysis, values were grouped by inflation

level and session e.g. RV S1 for all volunteers was compared to RV S2 for all

volunteers. Additionally, to assess repeatability in the image domain voxel-wise correlation (25) was carried out where each of the six same-inflation inter-session image pairs were spatially aligned via deformable image registration (3), in order to facilitate computation of Spearman correlation coefficients as previously described (30). The repeatability limit was calculated as $1.96 \times \sqrt{2}s_w$, where s_w is the withinsubjects standard deviation calculated using SPSS (version 23, IBM) (21). Spearman's correlation was also used to assess the relationship between TCV and %VV and H_{score} along with the relationship between TCV and the absolute change of %VV and H_{score} over the two imaging sessions. Finally, a two-way repeated measures analysis of variance was performed to statistically validate the effect of lung volume on H_{score} and %VV where within subject factors were defined as the imaging session and lung inflation level, and multiple comparisons were carried out using the Tukey correction. Voxel-wise correlation and two-way repeated measures analysis of variance was not carried out for the ¹²⁹Xe data due to the reduced number of subjects scanned. **Results** Comparison of HP ¹²⁹Xe and HP ³He MRI at different inflation levels The SNR of the ¹²⁹Xe images was lower than the SNR of the ³He images, particularly at RV, RV+1L and FRC, as can be seen in Figure 3. The RV image of HV3 (129Xe, session 2) had complete loss of signal from posterior sections of the lung due to a coil sensitivity issue at the time of the experiment and was thus excluded from analysis. $^{129}\mbox{Xe}$ images had consistently lower %VV (p<0.0001) and higher H_{score} (p<0.0001) when compared to those obtained with HP ³He (Tables 3 and 4).

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248 The effect of lung inflation level on %VV and H_{score} The effect of lung inflation level on ³He and ¹²⁹Xe images acquired at different lung 249 volumes is shown in Figure 4 for volunteer 2. There was a trend towards increased 250 251 ventilation homogeneity at higher lung volumes, which was seen using both gases. For ³He data significant differences between H_{score} were found when comparing TLC 252 to all other lung volumes via the two-way analysis of variance (p<0.0001 for all). No 253 254 other significant differences in H_{score} between different inflation levels were found. 255 256 %VV also varied with lung volume as can be seen from the mean values of %VV and H_{score} shown in Table 4 which are visualized in Figure 5. For ³He data, %VV at RV 257 258 and FRC+1L were the only volumes that were significantly different from each other 259 when compared using the two-way analysis of variance (p=0.0155). Lung volume had 260 a significant effect on both %VV (p=0.0265) and H_{score} (p<0.0001). 261 When considering the ¹H MRI acquired in the same breath as ³He MRI, TCV 262 generated from the ¹H images correlated strongly with H_{score} (r=-0.75, p<0.0001) but 263 264 not with %VV (r=0.27, p=0.15). TCV had a weak correlation with the absolute change in %VV (r=-0.39, p=0.03) but not H_{score} (r=0.01, p=0.53). For the ¹H MRI 265 acquired in a separate breath to the ¹²⁹Xe MRI, TCV had a strong correlation with 266 H_{score} (r=-0.90, p<0.0001) and a moderate correlation with the absolute change of 267 268 H_{score} over the two sessions (r=-0.66, p=0.01). TCV had no significant correlation with %VV or the absolute change in %VV over both sessions (r=0.44, p=0.12 and r=-269 270 0.33, p=0.25 respectively). 271

Regardless of the acquisition volume increased H_{score} was seen in the posterior region of the lung (Figure 6) with the most posterior slice having a mean±SD H_{score} over all volunteers and inflation levels of 15.4±7.1% whilst all other slices combined had values of 9.8±3.1% when considering ³He data. Additionally, significant differences between the most posterior slice and the remaining slices (Table 5) of the image were seen at RV+1L and FRC+1L (p=0.0087 and p=0.031 respectively) whilst no significant difference was seen at RV, FRC and TLC (p = 0.1562, p=0.3125 and p=0.0790 respectively). Repeatability of %VV and H_{score} Table 6 shows the CoV of %VV and H_{score} over all 6 volunteers at each of the lung volumes imaged with ³He and over all 3 volunteers imaged with ¹²⁹Xe. For ³He data, CoV was less than 1.5% for %VV and less than 5.5% for H_{score} at all lung volumes. Concerning 129 Xe data, CoV was less than 4% for %VV and less than 10% for H_{score} at all lung volumes. Concerning the ³He data, strong inter-session voxel-wise correlation was observed for all lung volumes (mean±SD Spearman coefficients: 0.92±0.03 for RV; 0.94±0.03 for RV+1L; 0.95±0.02 for FRC; 0.95±0.03 for FRC+1L; 0.93±0.02 for TLC). Bland-Altman bias±limits of agreement (LOA) are visualized in Figures 7 (³He) and 8 (129Xe) for both %VV (A) and H_{score} (B). For ³He data, the limits of agreement were less than 5% for %VV, and less than 2.5% for H_{score}. For ¹²⁹Xe data, the limits of agreement were less than 10% for %VV, and less than 4% for H_{score}. For ³He MRI the bias for %VV was less than 2% at all lung volumes whilst H_{score} bias was less than

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1% at all lung volumes whilst for 129 Xe MRI %VV bias was less than 6% at all lung volumes and H_{score} bias was less than 2% at all lung volumes.

Table 7 details the repeatability limit for %VV and H_{score} from both HP 3 He and 129 Xe images. When considering 3 He data %VV repeatability was less than 3% for all volumes except RV and less than 2% for all volumes when considering H_{score} . When considering 129 Xe data %VV repeatability was less than 10% for all volumes except RV and less than 3% for all volumes when considering H_{score} .

Discussion

The work carried out here has demonstrated that lung volume has a significant bearing on quantitative measurements of lung ventilation derived from both ³He and ¹²⁹Xe MRI. Additionally, from the effect of lung volume on the quantitative metrics of %VV and H_{score} evident in healthy volunteers, it can be concluded that the lung volume during imaging must be well controlled to ensure that these metrics can be used reliably in longitudinal studies.

Imaging over all volunteers revealed increased ventilation heterogeneity at lower lung volumes, potentially indicating partial airway closure in certain regions of the lung. Increased heterogeneity was particularly observed in the posterior section of the lung at RV+1L, exemplified by the median H_{score} per slice plotted against slice number for V5 in Figure 9. This increased heterogeneity is likely due to the breathing maneuver used to obtain the images at RV+1L, that is the volunteers first exhaled to RV, which may have caused some airway closure. In contrast, the HP gas mixture was inhaled from FRC for all other lung volumes, and so the ventilation seen in the RV and FRC

images was influenced by the gas distribution within the lungs at FRC+1L. Note that although increased heterogeneity is seen in the anterior portion of the lung, the increased H_{score} in those areas are due to the reduced SNR due to decreased gas reaching those areas within the lung.

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This increased ventilation heterogeneity at RV+1L in volunteers suggests the same underpinning mechanisms as reported in the work by Muradyan et al. (23), where there were distinct focal areas of lung affected by airway closure after inhalation of small gas volumes from below residual volume in elite divers. We hypothesize that the areas of decreased ventilation signal at RV+1L were caused by airways remaining closed following inhalation of the gas mixture. We believe that this same effect was not observed at RV in the current study since the maneuver to RV required first inhaling to FRC+1L, such that gas would remain in the areas opened by this first inhalation maneuver even if the airways were to close later on. The areas of reduced ventilation in lungs of the elite divers following inhalation from sub-RV levels observed in the work by Muradyan et al. (23) were larger than those seen here in these volunteers, whilst they did not see the same heterogeneity seen here in their volunteers following inhalation from RV. One possible reason for this is the improvements in the image resolution for ¹²⁹Xe when compared to their experiments that were carried out with 2D projection imaging, and thus providing us with better spatial sampling of regional heterogeneity.

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Imaging after smaller inspirations from RV would be interesting in order to assess at which point the ventilation heterogeneity would return to a distribution closer to that seen at FRC or FRC+1L. In this case, it would be expected that the smaller the

volume inhaled from RV, the greater the ventilation heterogeneity would be; although the feasibility of these experiments would be limited by the volume of HP gas required for sufficient image SNR if carrying the experiment out with 129 Xe. Another factor which may contribute to increased H_{score} at RV when compared to FRC+1L and FRC is the increased ratio of blood vessel volume to lung volume at RV, resulting in increased H_{score} .

The small CoV of %VV between sessions further confirms the growing body of evidence that %VV is a robust global metric of lung ventilation (5, 12, 17), and the high inter-scan repeatability makes %VV (or VDP) a good candidate metric for longitudinal assessment of lung function in patients (17). The proportionally larger CoV of the H_{score} suggests that this measure of global ventilation image heterogeneity may be less repeatable.

The generally lower SNR of ¹²⁹Xe images when compared to ³He images is a well-known phenomenon and follows previous publications (14, 28), with ¹²⁹Xe acquisitions having a mean±SD SNR of 30±13 compared to the 42±15 of the ³He acquisitions. Consequently, the higher H_{score} seen in the ¹²⁹Xe images when compared to images acquired with HP ³He is at least partially due to the lower SNR and thus increased heterogeneity of signal within ventilated regions. The lower %VV values measured from ¹²⁹Xe images compared to ³He images may be due to the lower diffusivity of ¹²⁹Xe compared to ³He, and are consistent with %VV values reported previously in healthy volunteers, patients with chronic obstructive pulmonary disease and patients with lung cancer who were imaged with both gases (18, 27). Furthermore, lower SNR in one of the ¹²⁹Xe acquisitions (V6, RV, S1) caused an increase in H_{score}

showing that the maneuvers or gas doses need to be optimized for the ¹²⁹Xe imaging acquisitions if this methodology is applied to patient cohorts. The need to register the anatomical images to the ventilation images for ¹²⁹Xe %VV calculation will also contribute to the lower repeatability of ¹²⁹Xe %VV when compared to ³He %VV (12), where anatomical images were acquired in the same breath-hold. Additionally, due to imaging constraints, HP ¹²⁹Xe images were acquired with double the slice thickness (10mm) of the HP ³He images (5mm); thus, differences would be expected due to different inherent physical properties and image acquisition considerations of the respective gas.

Imaging patients with HP gas at different lung volumes may provide a clearer picture of the nature of lung disease. For example, in patients with obstructive lung disease, following deep inhalation to TLC, the effect of increased positive pressure within the airways may result in a reduced H_{score} and increased %VV due to opening of obstructed airways (20). Additionally, as patients with chronic respiratory disease may have increased closing volumes, imaging at expiration may identify areas of gas trapping similar to those observed by Holmes et al. (9-11).

An increased number of healthy volunteers with a larger age range, and inclusion of female subjects would extend this preliminary work into the effect of lung volume on ventilation heterogeneity in healthy volunteers. Additionally, mitigating the signal dropout seen in TLC images with larger coil coverage is an important consideration for future studies. The smoking history of four of the six volunteers (two former smokers and two occasional smokers) means that these data may not represent the ventilation patterns seen in a group of healthy never-smokers. However, the number

of pack years reported by the volunteers scanned was low (<0.7), and in a previous ³He MRI study of pulmonary ventilation (26) three of the smokers would have been classified as never-smokers (<0.5 pack years). However, the volunteers scanned were spirometrically defined as free from respiratory disease and not unrepresentative of the general population in terms of smoking history. The fact that increased ventilation heterogeneity at lower lung inflation levels was seen in the two never smokers as well as those with a smoking history suggests this effect is not due to smoking related obstructive airways disease.

Conclusions

Increased ventilation heterogeneity was observed in HP gas images acquired at lower lung volumes in healthy volunteers. This work has shown that although TLV and VV may vary considerably between repeated scans there was little effect on %VV in these healthy volunteers. This indicates it may be important to image patients over a range of lung volumes with different breathing maneuvers to fully understand disease progression and accurately characterize ventilation defects and pulmonary mechanics. Finally, the variation in lung volume must be considered when monitoring patients longitudinally with hyperpolarized gas MRI particularly in the cases of disease with a reversible nature such as asthma.

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555 FIGURE LEGENDS 556 Figure 1 Example of local H_{score} calculation in the RV+1L ventilation-weighted image of volunteer 5 (top image). The yellow box on the left shows an area of low 557 H_{score} (enhanced image on the left, with the local area outlined and the voxel that is 558 559 replaced denoted with an 'x'), which is highlighted with the blue box in the H_{score} map (lower image). The same is shown for a region of high H_{score} on the right. 560 561 Figure 2 Breathing maneuvers and acquisition volumes used in this study. Solid gray 562 lines indicate an inhalation from a 1L bag, solid black lines indicate an exhalation and 563 dashed gray lines indicate an inhalation of room air. Solid boxes represent acquisition 564 volumes and dashed boxes represent intermediate volumes as part of the breathing 565 maneuver. Figure 3 Signal-to-noise ratio (SNR) values from the volunteers scanned with both 566 ³He and ¹²⁹Xe only. 567 Figure 4 Representative slices from all acquisition volumes in V2 from both ³He and 568 ¹²⁹Xe images. The top row shows the ³He images and the bottom row the ¹²⁹Xe 569 570 images acquired in V2. Figure 5 Plots of (A) percentage lung ventilated volume from ³He data, (B) H_{score} (%) 571 from ³He data, (C) percentage lung ventilated volume from ¹²⁹Xe data and (D) H_{score} 572 (%) from ¹²⁹Xe data at each acquisition volume. Each circle represents a volunteer 573 574 whilst the lines represent the mean of the values.

Figure 6 Representative posterior slices of HP ³He ventilation images and

heterogeneity maps at all acquisition volumes from V2. The arrows are pointing to

areas of decreased ventilation and increased H_{score}.

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578	Figure 7 Bland-Altman plots of (A) %VV and (B) H _{score} generated from images
579	acquired with HP ³ He at all acquisition volumes. Black dots indicate bias, gray dots
580	are the 95% confidence intervals and the black dashed line is 0.
581	Figure 8 Bland-Altman plots of (A) %VV and (B) H _{score} generated from images
582	acquired with HP ¹²⁹ Xe at all acquisition volumes. Black dots indicate bias, gray dots
583	are the 95% confidence intervals and the black dashed line is 0.
584	Figure 9 Exemplary plot of H _{score} from anterior to posterior for V5.
585	

Table 1 Subject demographics. V = volunteer, $FEV_1 = Forced$ expiratory volume in 1 second % predicted

Subject	Age, yr	Height, cm	Weight, kg	FEV ₁	Pack years
V1	32	183.0	87.0	102.0	0.15
V2	35	184.0	76.0	77.2	0.13
V3	31	182.0	83.0	105.0	0.06
V4	34	185.6	94.0	83.6	0.70
V5	27	189.5	74.0	102.9	0
V6	28	187.6	90.0	99.9	0

Table 2 Gas doses for hyperpolarized (HP) helium-3 (3 He) and xenon-129 (129 Xe) acquisitions reported as HP gas dose (N_{2}), where N_{2} = nitrogen. RV = residual volume, RV+1L = residual volume plus 1 liter of gas mixture, FRC = functional residual capacity, FRC+1L = functional residual capacity volume plus 1 liter of gas mixture and TLC = total lung capacity

Acquisition	³ He (N ₂), ml	129 Xe (N ₂), ml
RV	200 (800)	1000 (0)
RV+1L	150 (850)	750 (250)
FRC	200 (800)	1000 (0)
FRC+1L	150 (850)	600 (400)
TLC	200 (800)	750 (250)

Table 3 Mean percentage lung ventilated volume (%VV) and median score of the heterogeneity map (H_{score}) values at each lung volume and session (session 1 (S1)/session 2 (S2)) over all volunteers derived from hyperpolarized helium-3 (3 He) and xenon-129 (129 Xe)

	RV S1	RV S2	RV+1L S1	RV+1L S2	FRC S1	FRC S2	FRC+1L S1	FRC+1L S2	TLC S1	TLC S2
%VV ³ He	95.65	97.17	97.39	97.84	97.30	97.80	98.18	98.05	97.33	97.98
H _{score} ³ He	10.47	9.98	10.12	10.1	9.37	9.23	9.10	9.20	7.55	7.39
%VV ¹²⁹ Xe	82.94	87.43	92.36	90.86	93.53	94.99	92.99	96.55	95.83	94.98
$H_{score}^{129}Xe$	15.89	14.92	11.51	12.29	10.83	10.99	11.09	10.53	8.71	8.24

Table 4 Average percentage lung ventilated volume (%VV) and median score of the heterogeneity map (H_{score}) values over session 1 and session 2 generated from hyperpolarized helium-3 (3 He) and xenon-129 (129 Xe) images for the three volunteers (V) scanned with both gases

Acquisition	%VV ³ He V2	% VV ¹²⁹ Xe V2	%VV ³ He V3	% VV ¹²⁹ Xe V3	%VV ³ He V6	%VV ¹²⁹ Xe V6
RV	98.68	96.26	97.03	NA	95.41	88.88
RV+1L	97.67	90.46	99.49	91.54	98.26	92.84
FRC	98.85	97.53	98.34	94.41	97.85	90.85
FRC+1L	98.46	96.70	98.76	94.51	98.10	93.11
TLC	99.46	95.58	99.12	93.14	97.82	97.50
Acquisition	H _{score} ³ He V2	H _{score} ¹²⁹ Xe V2	H _{score} ³ He V3	H _{score} ¹²⁹ Xe V3	H _{score} ³ He V6	H _{score} ¹²⁹ Xe V6
RV	8.96	12.53	11.63	NA	10.16	16.16
RV+1L	10.26	12.08	8.82	11.74	9.04	11.89
FRC	8.85	11.01	9.15	10.91	8.90	10.82
FRC+1L	9.33	10.83	8.61	10.62	8.59	10.98
TLC	6.15	8.10	7.33	8.95	6.50	8.38

Table 5 Mean H_{score} (over session 1 and session 2) at the most posterior slice and all remaining slices for all volunteers at each lung volume for all images acquired using hyperpolarized helium-3

	RV		RV	'+1L	FR	RC	FRC+1L		TLC	
	Posterior	Remaining								
Volunteer	slice	slices								
V1	14.95	11.75	21.38	12.11	23.9	12.19	21.13	10.72	17.45	10.3
V2	12.68	9.13	23.97	11.62	15.11	9.30	23.59	10.22	9.28	6.85
V3	8.28	11.15	18.18	8.58	5.59	9.06	10.16	8.65	7.64	7.40
V4	20.23	10.91	17.83	10.24	18.15	10.65	20.46	10.74	25.71	9.39
V5	9.98	10.98	27.03	11.03	5.37	9.42	9.53	9.60	8.29	8.25
V6	15.57	11.00	9.58	9.57	11.53	9.40	13.60	9.02	27.03	7.18

Table 6 Coefficient of variation (CoV) at each inflation level for metrics derived from hyperpolarized helium-3 (³He) and xenon-129 (¹²⁹Xe)

		³ He			¹²⁹ Xe				
Acquisition	TLV	VV	%VV	H _{score}	Acquisition	TLV	VV	%VV	H _{score}
RV	3.40	3.05	1.29	5.32	RV	3.33	1.34	3.98	9.37
RV+1L	4.13	4.64	0.63	4.62	RV+1L	2.19	2.26	1.16	7.60
FRC	4.63	4.64	0.87	3.99	FRC	5.88	4.80	1.49	2.86
FRC+1L	3.42	3.42	0.38	2.74	FRC+1L	6.88	6.00	3.18	3.74
TLC	1.19	1.00	0.54	5.46	TLC	1.97	1.91	0.62	4.62

Table 7 Repeatability limit for percentage lung ventilated volume (%VV) and median value of the heterogeneity map (H_{score}) for images acquired using hyperpolarized helium-3 (^{3}He) and xenon-129 (^{129}Xe)

Acquisition	³ He %VV	³ He H _{score}	¹²⁹ Xe %VV	¹²⁹ Xe H _{score}
RV	5.08	1.80	11.69	3.72
RV+1L	2.19	1.86	3.06	2.62
FRC	2.90	1.29	4.50	1.07
FRC+1L	1.39	0.80	9.59	1.45
TLC	2.02	1.35	1.95	1.36

















