Lung clearance index is a sensitive, repeatable and practical measure of airways disease in adults with cystic fibrosis

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ABSTRACT

Background: Lung clearance index (LCI) is a sensitive marker of early lung disease in children but has not been assessed in adults. Measurement is hindered by the complexity of the equipment required. The aims of this study were to assess performance of a novel gas analyser (Innocor) and to use it as a clinical tool for the measurement of LCI in cystic fibrosis (CF).

Methods: LCI was measured in 48 healthy adults, 12 healthy school-age children and 33 adults with CF by performing an inert gas washout from 0.2% sulfur hexafluoride (SF6). SF6 signal:noise ratio and 10–90% rise time of Innocor were compared with a mass spectrometer used in similar studies in children.

Results: Compared with the mass spectrometer, Innocor had a superior signal:noise ratio but a slower rise time (150 ms vs 60 ms) which may limit its use in very young children. Mean (SD) LCI in healthy adults was significantly different from that in patients with CF: 6.7 (0.4) vs 13.1 (3.8), p<0.001. Ten of the patients with CF had forced expiratory volume in 1 s ≥80% predicted but only one had a normal LCI. LCI reproducibility in all three groups of subjects (mean intra-visit coefficient of variation ranged from 3.6% to 5.4%).

Conclusions: Innocor can be adapted to measure LCI and affords a simpler alternative to a mass spectrometer. LCI is raised in adults with CF with normal spirometry, and may prove to be a more sensitive marker of the effects of treatment in this group.

As part of a programme aimed at measuring the response to gene therapy in cystic fibrosis (CF), we are interested in developing more sensitive measures of changes in CF airway function and structure. In the USA the only marker of lung function currently recognised as a primary end point in CF trials is the forced expiratory volume in 1 s (FEV1).1 In early disease this reflects total airways resistance and is insensitive to changes in small airways, which contribute <10% of the overall resistance in healthy adult subjects.2 Significant structural airway damage can be demonstrated on CT scanning in the presence of a normal FEV1.3

In early lung disease, ventilation heterogeneity results from regional differences in small airway calibre (those beyond division 8).4 5 This can be demonstrated both in computer models of the human lung6 and from in vivo MR images7 or radiolabelled tracer gas distribution.8 Inert gas washout is an alternative technique which involves measuring the elimination of a non-absorbed gas as it is exhaled during tidal breathing. The gas can either be resident nitrogen washed out by breathing 100% oxygen, or an exogenous tracer gas which has first been breathed in to equilibrium. Lung clearance index (LCI), a simple marker of deranged ventilation, can be calculated from the washout curves.9 Past studies using a variety of methods have shown that LCI is reproducible and more sensitive than FEV1 to identifying early lung disease in children.2 10 In addition, Aurora et al10 showed that LCI is further raised in children infected with Pseudomonas aeruginosa and Kraemer et al11 showed LCI to be an early predictor of deteriorating lung function in children.

Although an old technique,10 measurement of LCI has always relied upon complex and bulky equipment, usually assembled from separate components by the investigators themselves, and has largely been restricted to a research setting. The current best method involves using a mass spectrometer (MS) to measure the washout of the inert tracer gas sulphur hexafluoride (SF6). Although these are simpler than nitrogen washouts, MS are expensive to purchase and maintain.

The purpose of this study was to investigate LCI in healthy subjects and adults with CF using a modified Innocor device (Innovision, Odense, Denmark). Innocor is a compact gas analyser and flow sensor originally designed to measure cardiac output by inert gas rebreathing. The gas analyser uses photoacoustic spectroscopy to measure several gases including low concentrations of the inert tracer SF6, making it a suitable device for ventilation distribution measurements. More information on the gas analyser is given in the online supplement (pages 2–5).

In preparation for its use as an end point in clinical trials, the aims of this study were:

1. To compare the performance (assessed by response time and signal:noise ratio) of the Innocor gas analyser with that of the current inert gas washout standard (MS).

2. To adapt the Innocor device and analysis software into a clinical system for measurement of functional residual capacity (FRC) and LCI.

3. To assess how LCI changes with age of subject in healthy volunteers.

4. To assess the intra and inter-visit reproducibility of LCI in healthy volunteers.

5. To use the adapted Innocor to measure FRC and LCI in normal adult subjects and patients with CF and to compare LCI with spirometry.
Cystic fibrosis

Figure 1 Patient interface used for inert gas washout with Innocor gas analyser.

METHODS

Equipment

To measure LCI, a mouthpiece fitted with a flowmeter and gas sampling port is required (fig 1). This is connected to a detachable flowpast tube which is used to supply tracer gas during the wash-in and is then removed at the start of washout. A more detailed Methods section is available in the online supplement, and the modifications to the standard Innocor patient interface are described in detail on pages 5–7.

Spirometry was measured according to American Thoracic Society/European Respiratory Society guidelines; predicted values for FEV₁ are those provided by the European Community for Coal and Steel (adults ≥17 years) and Rosenthal et al (children <16 years).¹⁷

Performance of Innocor gas analyser

The signal:noise ratio at the start and end of a washout and the rise time of the gas analyser in response to a step change in SF₆ concentration were assessed as described in the online supplement (page 8). Performance was compared with that of a MS used routinely for LCI measurements. The ability of the complete modified system to integrate flow and gas signals accurately was assessed using a gas calibration syringe which can be set to deliver different volumes (Hans Rudolph, Missouri, USA). This was filled with 0.2% SF₆ in air (BOC, Guildford, UK) to a range of different starting volumes and a washout performed by incomplete filling and emptying of the syringe around this starting point. The syringe volume derived from the calculated “expired” volume of SF₆ was then compared with the known starting volume.

Flow and SF₆ data were exported for analysis on custom-built software. FRC was derived from the total expired SF₆ volume, calculated by integration of flow and SF₆ signals. LCI is defined as the number of lung turnovers (ie, multiples of FRC) required to reduce end tidal marker gas concentration to 1/40th of the starting concentration (ie, <0.005%). The flowpast circuit was then detached during expiration and the washout measured until the end tidal SF₆ had fallen to less than 1/40th of the starting concentration (ie, <0.005%). In healthy children (<16 years) an identical gas analyser and protocol were employed at a separate research site, but a smaller filter was used to reduce the precapillary dead space (36 ml vs 46 ml in adults).

Subjects completed three sets of wash-ins and washouts. A washout was discarded if the resulting calculated FRC differed by >10% from both the other two repeats.¹⁵

Statistical analysis

Data were analysed using Prism (GraphPad Software Inc, California, USA). The results are given as mean (SD) unless otherwise stated. Within-test repeatability for LCI was determined by calculating the coefficient of variation (CV) as 100 × residual standard deviations. A p value of <0.05 was considered statistically significant.

RESULTS

Technical validation of Innocor device

Signal:noise ratio of Innocor and MS

The Innocor device has a lower gas concentration operating range than the MS. Signal quality is therefore given at the starting and finishing concentrations of a washout, which are different for the two devices (table 1). For both devices there is a fall in signal:noise ratio as the gas concentration falls, but the Innocor signal quality remains superior throughout, despite much lower SF₆ concentrations.

Rise time and delay of gas signal

The mean (SD) SF₆ 10–90% rise time was 154 (5) ms for Innocor and 64 (5) ms for the MS (p<0.001). The longer rise time of the Innocor gas analyser was allowed for by offsetting gas and flow signals during analysis by an additional 50 ms. This corresponds to the 50–80% rise time of the gas signal, and has the effect of speeding the response time by realigning the flow signal with the 80% response fraction of the gas signal.¹⁶

Subjects

Forty-nine healthy non-smokers (<10 pack-years smoking history) with no active lung disease and on no regular respiratory medications were recruited as normal adult volunteers (age range 19–58 years). Thirteen healthy child volunteers (age range 6–16 years) were recruited if they had no previous diagnosis of recurrent wheeze or asthma and were taking no current inhaled medication. There was no history of significant respiratory disease requiring hospitalisation (eg, pneumonia, pertussis, tuberculosis), no prematurity (<34 weeks gestation) and no significant co-morbidity. Thirty-three patients with CF (age range 17–49 years) were recruited from the Scottish Adult CF Service, the diagnosis being based on a combination of clinical presentation and sweat testing and confirmed by genotyping. All volunteers, patients and (where relevant) parents provided informed consent. Paediatric volunteers provided assent where appropriate. This study was approved by the Lothian research and ethics committee.

Washout test

Subjects were seated and suitably distracted by watching television. A noseclip was applied and tidal breathing established while the subject was connected to the flowpast circuit containing 0.2% SF₆ in air. This was supplied from a compressed gas cylinder with the flow rate adjusted to ensure that rebreathing did not occur. This wash-in phase continued for at least 5 min in adults or 4 min in children under 16 years and, in all cases, until inspiratory and expiratory SF₆ concentrations differed by <0.004% (absolute difference in SF₆ concentration). The flowpast circuit was then detached during expiration and the washout measured until the end tidal SF₆ had fallen to less than 1/40th of the starting concentration (ie, <0.005%). In healthy children (<16 years) an identical gas analyser and protocol were employed at a separate research site, but a smaller filter was used to reduce the precapillary dead space (36 ml vs 46 ml in adults).

Subjects completed three sets of wash-ins and washouts. A washout was discarded if the resulting calculated FRC differed by >10% from both the other two repeats.¹⁵
The accuracy of this adjustment was then confirmed by integration of known volumes of SF₆ from a calibration syringe.

Validation of FRC measurements
Sixteen washouts were performed using a calibration syringe with the starting volume varied between 1.5 and 3 litres. There was good agreement between the measured and actual syringe volumes (see fig 1 in online supplement II). The mean (SE) error between measured and actual syringe volume was 16.3 (2.4) ml or 1.1 (0.2)%.

In vivo LCI measurement
LCI was assessed successfully in 12 healthy children, 48 adult healthy volunteers and 33 adults with CF. The demographic data of the study subjects are given in table 2. Data from two additional healthy volunteers (one adult, one child) could not be analysed because of technical difficulties (see below).

Effect of age, height and gender on LCI in non-CF subjects
Figure 2 shows the relationship between LCI and age (min 6 years, max 58 years). In those aged >16 years there was no relationship between LCI and age. When the two cohorts were combined there was a weak but statistically significant correlation with age (Pearson r² = 0.16, p<0.002). The small dependence of LCI on age is best summarised by a normal range (95% limits of normality) in adults of 5.9 to 7.5 and in children (<16 years) of 5.3 to 7.3. A weak relationship between height and LCI in the combined cohorts disappeared on multiple regression analysis. LCI was unrelated to gender of subject. By contrast, FEV₁ varied between 76% and 133% predicted in the combined cohorts.

LCI in non-CF and CF adults
The group mean (SD) LCI in adult healthy controls was 6.7 (0.4) (range 6.0–7.8) with 95% limits of normality calculated as 5.9 to 7.5. In patients with CF the group mean (SD) LCI was 12.8 (5.6) (range 6.3–20.4), p<0.001 compared with healthy controls. The mean (SD) FEV₁ was also significantly different between the two groups (102 (12)% predicted in healthy controls vs 68 (25)% predicted in patients with CF, p<0.001).

Effect of age, height and gender on LCI for healthy controls and adults with CF. In controls LCI was restricted to a narrow range but, in patients with CF, LCI increased with reducing FEV₁ % predicted (r² = 0.69, p<0.001).

There were 10 patients with CF with FEV₁ ≥80% predicted, all but one of whom had LCI above the upper limit of normal. By contrast, LCI was marginally raised in only two healthy adults (measuring 7.7 and 7.8). The sensitivity of LCI for detecting CF was 97% compared with 70% for FEV₁.

Repeatability of washout at same visit
A washout test was excluded if the measured FRC differed by >10% from both of the other two washouts. In adult subjects this resulted in the exclusion of a total of seven tests, representing <3% of the total number of repeats from both healthy volunteers and patients with CF. Three tests were excluded from the paediatric cohort, representing 9% of the total number of washout repeats. All three washout repeats from an additional single adult healthy volunteer could not be analysed because they were unable to achieve a regular and reproducible breathing pattern. All three repeats from an additional healthy child (age 8) were also excluded because of evidence of an air leak.

After exclusion of these repeats, the mean (SD) intra-subject coefficient of variation (CV) for FRC derived from repeat washout manoeuvres on the same visit was 3.2 (1.9)% for adult healthy volunteers, 3.9 (2.1)% for healthy children and 3.5 (2.5)% for patients with CF. The mean (SD) CV for LCI was 3.6 (2.1)% for healthy adults, 5.4 (3.8)% for healthy children and 4.4 (2.8)% for patients with CF. There was no significant correlation between the LCI CV and FEV₁ % predicted.

Inter-visit reproducibility of LCI in healthy adults
Repeat measurements of LCI were performed in triplicate on 16 healthy volunteers after a mean (SE) of 56 (10) days.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Signal/noise ratios of Innocor and mass spectrometer (MS) at gas concentrations encountered at start and end of washout</th>
</tr>
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<tbody>
<tr>
<td>SF₆ concentration (%)</td>
<td>MS</td>
</tr>
<tr>
<td>Start</td>
<td>End</td>
</tr>
<tr>
<td>4.0</td>
<td>0.1</td>
</tr>
<tr>
<td>0.2</td>
<td>0.005</td>
</tr>
</tbody>
</table>

The signal/noise ratio is calculated as the ratio of mean to standard deviation of a stable gas signal over 10 s.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Demographic data, spirometric parameters and lung clearance index (LCI) of healthy volunteers and patients with cystic fibrosis (CF)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Healthy volunteers</td>
<td>Children (age &lt;16 years)</td>
</tr>
<tr>
<td>N</td>
<td>12</td>
</tr>
<tr>
<td>M/F</td>
<td>7/5</td>
</tr>
<tr>
<td>Mean (range) age (years)</td>
<td>11 (6–16)</td>
</tr>
<tr>
<td>Mean (SD) FEV₁ (% predicted) LCI</td>
<td>95 (11)</td>
</tr>
<tr>
<td>Mean (SD)</td>
<td>6.3 (0.5)</td>
</tr>
<tr>
<td>Range</td>
<td>5.6–7.1</td>
</tr>
<tr>
<td>Mean (SD) CV%</td>
<td>5.4 (3.8)</td>
</tr>
</tbody>
</table>

FEV₁, forced expiratory volume in 1 s; CV%, coefficient of variation (%) for intra-visit repeats.

*p<0.001 vs adult healthy volunteers (Mann-Whitney U test).
A Bland-Altman plot of the difference between repeat measures and the mean of the measurements for LCI is shown in fig 4. For FRC, the 95% limits of agreement between the two measurements were $-0.43$ to $0.45$ litres and, for LCI, the 95% limits of agreement for the two measurements were $-0.78$ to $0.46$. The inter-visit reproducibility of the FRC measurement was therefore approximately 400 ml and that of the LCI measurement was 0.6.

**DISCUSSION**

This study has shown that the clinical measurement of inert gas washout is practical using equipment that is cheaper, more portable and has more sensitive gas signal resolution than the current MS standard. We have also shown for the first time that, in adults with CF, a simple measure of ventilation heterogeneity is more sensitive than FEV$_1$ in detecting lung function abnormalities. Finally, we have shown that this measurement is reproducible both within and between visits, and that there is little change over a wide range of subject height and age.

In children with CF there is already increasing evidence that LCI is a more sensitive measure of early lung disease than FEV$_1$. LCI has also been shown to correlate better with scores of airway damage on high-resolution CT scanning than spirometry. It may therefore fill an important gap in our ability to monitor airway function and disease progression non-invasively. Since only tidal breathing is required, it is particularly suitable for airways assessment in subjects who find complex respiratory manoeuvres difficult.

The potential of multiple breath washout measurements, however, has been hampered by the lack of a convenient method of performing them. The original method for assessing lung clearance was the nitrogen washout. Although this avoids the need for a wash-in first, sufficient time must be allowed between tests for the end tidal nitrogen concentration to return to normal. The use of a MS to measure LCI by following changes in exogenous SF$_6$ is now well described in children and is probably the accepted gold standard technique in this population. The MS offers the additional advantage that it can measure a wider range of different gases, which is a useful option when measuring vital capacity single breath washouts. However, the MS is an expensive, temperamental and bulky piece of equipment that cannot readily be taken out of the laboratory. In contrast, Innocor contains both the gas analyser and the pneumotachograph in a single unit that is both portable and robust. A supply of SF$_6$ is required for both systems, but the concentration required for Innocor is 1/20th of that used in the MS washouts, which reduces gas wastage and the potential environmental (greenhouse) effects of SF$_6$.

The ideal comparison would be to compare the performance of both systems simultaneously, as has been done for other gas analysers. However, washouts would have to be performed at the operating range of the Innocor gas analyser since the response is not linear above 0.2% SF$_6$, but the signal resolution of the MS shows excessive noise at this level. Accepting this as a limitation of the current comparison, we have shown that the gas analyser is suitable for use in a multiple breath washout apparatus. The characteristics of the two analysers are summarised in table 2 of the online supplement (page 18). Our technical validation shows that the device with our modifications is capable of measuring gas volume by dilution with high accuracy. Despite operating at a much lower SF$_6$ concentration, it produces washouts with a superior signal:noise ratio than a MS. Our comparison has also identified the possible limitations of the device imposed by the slower signal response.

**Figure 2** Effect of age on lung clearance index (LCI) in healthy volunteers. LCI remains within a narrow band of normal over an age range of 52 years. The broken line is the regression line, showing the extent of age-related increase in LCI.

**Figure 3** Lung clearance index (LCI) versus forced expiratory volume in 1 s (FEV$_1$) % predicted for adult healthy volunteers and patients with cystic fibrosis. The horizontal line represents the mean and the horizontal dotted lines the 95% limits of normality of the LCI, calculated from the healthy adult population. The vertical line represents the lower limit of normal for % predicted FEV$_1$.

**Figure 4** Bland-Altman plot of difference between lung clearance index (LCI) measured on two separate occasions (quoted as mean of triplicate repeats) and mean of the two measurements of LCI.
rise and fall time. The system is able to integrate flow and \( \text{SF}_6 \) concentrations accurately at a physiological breathing rate of 10–30 breaths/min and should therefore be suitable for use in school-age children and adults. The response time may, however, limit the use of the method in neonates and preschool children with faster respiratory rates.\(^{21,27} \) Further in vitro assessment is required before using the analyser in this age group.

To date, we have used the modified Innocor to measure LCI in more than 100 patients and volunteers. From the data presented here, over 85% of subjects are able to complete all three washout manoeuvres without difficulty and generate reproducible measurements of FRC and LCI. Even for patients with CF, the whole process (wash-in and washout) usually takes little more than 10 min, and considerably less in children. Despite the relatively uncontrolled conditions, the mean CV for repeat FRC is similar to that described in the literature, which varies from 3.5% to 6.7% for plethysmography and from 4.9% to 10.4% for helium dilution.\(^{29} \) The mean CV for LCI is also better than that described in children.\(^{13} \) Repeat measurements of LCI at a separate time point in a cohort of adult healthy volunteers also demonstrated good inter-visit reproducibility.

It has been shown that LCI may be influenced by large changes in tidal volume, respiratory rate or FRC.\(^{29,30} \) We used tidal volume feedback to control tidal volume and respiratory rate within a range which should not affect the result. Since LCI is a ratio of cumulative expired volume and FRC, it is also independent of small changes in FRC over the physiological range. This is supported by the reproducibility of LCI and the narrow range of LCI in normal subjects found in the current study. Furthermore, because it is normalised for FRC, the normal range of LCI is largely unaffected by age, height or gender of subject. There was a weak but statistically significant rise in LCI with age. The clinical significance of this is unclear, since the magnitude of the difference (over a 52-year age range) remains very small and is less than inter-visit reproducibility. Serial dead space is known to affect LCI in infants and remains very small and is less than inter-visit reproducibility.\(^{11} \) However, the change in normal values of LCI was therefore possible that this represents a true effect of age on lung elasticity and hence gas mixing. By contrast, there is a wide range of “normal” for FEV\(_1\) % predicted, which is influenced by the choice and accuracy of the normal range selected.\(^{52} \)

The mean (SD) LCI determined here is very similar to that reported in the literature in children and adolescents. In preschool children (mean age 4 years) this has been reported as 6.9 (0.4),\(^{16} \) and in school-age children (mean age 11 years) as 6.5 (0.5)\(^{27} \) and 6.5 (0.4)\(^{16} \) in two different populations from the UK and Sweden, respectively. This supports our observation that LCI changes little with the age of the subject (>6 years). This may be especially useful during the long-term follow-up studies.

These are the first data on LCI in adults with CF; previous studies have only reported measurements in subjects up to 19 years of age. Even in adult patients with normal spirometry, the LCI may be markedly elevated, indicating significant “silent” lung damage. Some of the patients with normal FEV\(_1\) had no symptoms and were on no treatment, the diagnosis of CF having been made incidentally. Yet, despite this, there was abnormal gas mixing in almost all cases. There is a risk that the extent of lung disease in such patients will be underestimated and hence undertreated.

While FEV\(_1\) is the currently accepted gold standard to monitor trials of new treatments for CF, the rate of decline in this parameter has steadily reduced over the last decade.\(^{23} \) Power calculations show that many hundreds of patients would need to be treated over a year or more to see any beneficial effect of a novel therapeutic agent aimed at the basic defect.\(^{24} \) We have therefore instituted a large programme to assess novel biomarkers which could act as surrogates for FEV\(_1\). Ventilation heterogeneity is thought to be altered by small airways dysfunction.\(^{4,5} \) and measurements of this should therefore reflect the earliest pathology in CF—as has already been shown in children.\(^{11} \) This is also the region of the lungs which is likely to be a key target for gene therapy. The choice of which subject to recruit into trials of gene therapy represents a conflict between those with sufficiently clear airways that the gene therapy complex can be delivered into the lungs and those with sufficient abnormality in lung function so that any improvement can be measured. LCI offers the ability to measure dysfunction in the airways of interest, and also to extend the range of patients suitable for these assessments.

We have shown that there is the possibility of a robust and compact apparatus to measure LCI that can be used in multicentre studies after relatively straightforward modification. This will permit us to assess LCI routinely in patients to obtain longitudinal data and, in particular, it may serve as a more sensitive measure of lung function after changes in therapy. The value of this technology may, however, extend beyond just CF and we anticipate that it may provide valuable information about the development and treatment of airways disease in other conditions. In particular, it may be useful in conditions that initially affect the small airways such as asthma, chronic obstructive pulmonary disease and obliterative bronchiolitis.

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REFERENCES

Cystic fibrosis


Lung alert

Extracellular ATP in asthma airway inflammation

A role for ATP as a pro-asthmatic mediator has previously been suggested from in vitro studies. This group investigated whether extracellular ATP and purinergic signalling are important mediators of airway inflammation in asthma, using experimental mice models and human subjects.

Rapid accumulation of ATP was observed in bronchoalveolar lavage (BAL) fluid after allergen challenge in subjects with allergic asthma and in experimentally sensitised mice. In the mouse model, ATP binding to purinergic receptors drove inflammatory chemotaxis and bronchospasm. Prevention of this increase in ATP using treatment with the ATP-hydrolysing enzyme apyrase to degrade ATP, or blockade of ATP effects using broad spectrum purinergic P2 receptor antagonists, down modulated all of the cardinal features of the asthmatic response. In addition, administration of exogenous ATP to mice enhanced Th2-type sensitisation to inhaled antigen, indicating that ATP may contribute to the development of asthmatic sensitisation by recruitment of Th2-inducing lung myeloid dendritic cells to mediastinal lymph nodes.

The authors conclude that ATP and purinergic signalling are important in the pathogenesis of asthma. Further studies are needed, but this may be an important therapeutic target for the future.


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