

ORIGINAL RESEARCH

Lung function and asthma control in school-age children managed in UK primary care: a cohort study

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ABSTRACT

Background Spirometry and fraction of exhaled nitric oxide (FeNO) are commonly used in specialist centres to monitor children with asthma. The National Institute for Health and Care Excellence recommends spirometry for asthma monitoring from 5 years in all healthcare settings. There is little spirometry and FeNO data in children managed for asthma in UK primary care to support their use.

Objectives To study the prevalence of abnormal spirometry and FeNO in children with asthma managed in primary care and to explore their relationship with asthma control and unplanned healthcare attendances (UHA).

Methods Prospective observational cohort study in children aged 5–16 years with suspected or doctor-diagnosed asthma attending an asthma review in UK general practice. Spirometry, FeNO, asthma control test (ACT) scores and number of UHAs were studied.

Results Of 612 children from 10 general practices, 23.5% had abnormal spirometry, 36.0% had raised FeNO ≥ 35 parts per billion and 41.8% reported poor control. Fifty-four per cent of children reporting good asthma control had abnormal spirometry and/or raised FeNO. At follow-up, the mean number of UHAs fell from 0.31/child in the 6 months preceding review to 0.20/child over the 6 months following review ($p=0.0004$). Median ACT scores improved from 20 to 22 ($p=0.032$), and children's ACT from 21 to 23 ($p<0.0001$).

Conclusions Abnormal lung function and FeNO are common in children attending for asthma review in primary care and relate poorly to symptom scores. A symptoms-based approach to asthma monitoring without objective testing is likely to miss children at high risk of future severe asthma attacks.

Key messages

What is the key question?

- ▶ How common is abnormal spirometry and fraction of exhaled nitric oxide (FeNO) in children managed for asthma in primary care, and what is their relationship with patient-reported asthma control?

What is the bottom line?

- ▶ Abnormal spirometry and FeNO are prevalent in children cared for in primary care, and a symptoms-based asthma assessment alone will under-recognise children with obstructed airflow or active airway inflammation.

Why read on?

- ▶ To our knowledge, this is the first primary care study to explore the role of spirometry and FeNO testing in children managed for asthma in the UK.

control and 2) identifying risk factors for future poor outcomes, which include poor lung function.

The importance of assessing future asthma risk was also highlighted in the 2014 UK National Review of Asthma Deaths. This report found that the severity of asthma was underestimated in two-thirds of children who died from asthma between 2012 and 2013,³ and identified deficiencies in the recognition of patients at high risk of asthma attacks and death. One of the key recommendations from the report was that 'practices need to adopt a system of establishing the risk profile of a patient and put a treatment plan in place that is appropriate to their risk profile'.

GINA recommends that spirometry is recognised as an important part of the assessment of future asthma risk,⁴ and more recently the UK National Institute for Health and Care Excellence (NICE) published asthma guidelines⁵ recommending the use of spirometry for asthma monitoring in children aged 5 years and over.

The role of fraction of exhaled nitric oxide (FeNO) monitoring is less clear, although there is some evidence to suggest that FeNO-guided asthma treatment reduces exacerbation rates in children when compared with guidelines-based asthma management.⁶ GINA lists raised FeNO as a risk factor for future asthma attacks, at least in adults, while NICE recommends consideration of FeNO measurements to support asthma management in

INTRODUCTION

The UK has one of the highest asthma mortality rates among high-income countries in Europe in young people aged 10–24 years.¹

The existing UK British Thoracic Society/Scottish Intercollegiate Network Guideline defines good asthma control in terms of current symptoms control²; recommending that 'asthma is best monitored in primary care by routine clinical review', using standardised questionnaires but do not promote the need for objective lung function testing. This is in contrast with recommendations from the Global Initiative for Asthma (GINA), which divides assessment of asthma control into two domains: 1) evaluation of current symptom



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people who are symptomatic despite using inhaled corticosteroids. In primary care, spirometry and FeNO are rarely measured in the management of asthma in children or adults.

Implementation of spirometry and FeNO monitoring would represent a major change in routine primary care asthma management and has understandably raised concerns among stakeholders.⁷⁻⁹ This is chiefly because there is little UK data relating to lung function and asthma control in children managed in primary care.

We conducted a pragmatic prospective observational study in primary care with the objectives to quantify: 1) the prevalence of abnormal spirometry, FeNO and poor current asthma control at a single time-point in a cohort of children with either suspected or an existing diagnosis of asthma, and 2) the proportion of children in whom an assessment based on current symptoms alone would have failed to identify those with abnormal lung function and FeNO who may be at an increased risk of a severe asthma attack. As a secondary outcome, we explored the change in current asthma control and the number of unplanned healthcare attendances (UHAs) in the 6 months following a structured clinical review that included spirometry and FeNO testing.

METHODS

Prospective observational cohort study in children managed in primary care in the East Midlands, UK between 2016 and 2017.

Practices

Identification of general practices was facilitated by the Clinical Research Network. We sought to include practices of different sizes, serving populations of different ethnic and socioeconomic profiles. Seventeen general practices were contacted and 10 expressed an interest to participate.

Participants

Children aged 5–16 years with an existing asthma diagnosis or children receiving asthma medications (suspected asthma) were eligible to participate. Children fulfilled the inclusion criteria if at least one of the following was present:

1. On the practice asthma register.
2. Prescribed inhaled corticosteroids within the previous 12 months.
3. Prescribed ≥ 2 short acting beta-2-agonist via metered dose inhalers (SABA MDIs) in the previous 12 months.
4. Prescribed oral corticosteroids for acute wheeze/cough/breathlessness in the previous 12 months.

Eligible children were identified from the electronic register of each participating practice and invited by letter to attend the practice for an asthma review. Families who did not respond to the first letter were sent a second invitation and/or contacted by telephone by their registered practice.

Procedures

Training in performing and interpreting spirometry and FeNO was provided for clinical staff at each practice by the research team.

Clinical staff were trained to review and adjust treatment plans, and adherence to asthma preventer medications, of any child with either poor symptom control or abnormal objective test results. As all clinical reviews were conducted by the practice staff independent of the research team, the routine checking of inhaler technique and issuing of written personalised asthma action plans at each clinical review was encouraged but not monitored as part of this study.

Written consent was sought from all families attending for an asthma review for the purposes of data collection. Spirometry and FeNO testing were attempted in all children.

Asthma control tests

Validated questionnaires were used to assess current asthma control; the asthma control test (ACT)¹⁰ for children aged ≥ 12 years, and the childhood asthma control test (CACT)¹¹ for children aged 5–11 years. A score of ≤ 19 in either questionnaire denotes current uncontrolled asthma, with better control indicated by higher values.⁵

Spirometry

Spirometry was performed using a portable spirometer (Care-Fusion, UK) by practice staff either directly supervised or independently once competent. Forced expiratory manoeuvres were performed according to American Thoracic Society and European Respiratory Society (ATS/ERS) standards.¹²

Predicted values were based on global lung initiative (GLI) reference equations.¹³

For training purposes, a normal spirogram was defined as having both FEV₁ and FVC $\geq 80\%$ predicted and FEV₁/FVC $\geq 80\%$. A fixed cut-off was chosen to define obstruction for pragmatic reasons, as it was felt that general practice staff were already familiar with using fixed cut-offs when interpreting adult spirometry. The 80% threshold was chosen as it most closely mirrors the GLI lower limits of normal (LLN) across the 5–16 years age group.

Where FEV₁ or FVC was $< 80\%$ predicted, or FEV₁/FVC $< 80\%$, reversibility testing was performed: after administering salbutamol 400 μg via spacer. An increase in absolute FEV₁ (L) of $\geq 12\%$ was taken to represent a positive bronchodilator response (BDR) test.

Fraction of exhaled nitric oxide

FeNO measurements were attempted in all children following spirometry and performed in accordance with standardised guidelines.¹⁴ We used a handheld FeNO analyser (NIOX Vero; Circassia). We used a cut-off value of ≥ 35 parts per billion (ppb) to represent evidence of current active airway inflammation.⁵

Follow up

We reviewed each participant's electronic records 6 months following the initial asthma review. All record entries including consultations, electronically filed letters and prescriptions for the 6 months period were reviewed in order to obtain the number of UHA since the initial review. A UHA was defined as any attendance to the GP, hospital or walk-in centre with acute respiratory symptoms managed with asthma medications as documented within GP records or filed as an electronic discharge letter from hospital. In patients who consented, we also sent out follow-up letters with repeat ACT/CACT questionnaires.

Data analysis

We compared baseline characteristics, lung function test results and UHAs between children who were on the practice asthma register with those who were not. Continuous variables were compared using unpaired t-tests for parametric data, and Kruskal-Wallis and Wilcoxon signed rank tests for non-parametric data. Chi-squared tests were used for count data.

Lung function data were analysed according to the diagnostic status, age group and the current asthma symptom control status of the children, as binary variables. We used a fixed cut-off for

FEV₁ predicted and FEV₁/FVC of 80% to define airflow obstruction for training purposes. However for analysis, the gold standard GLI LLN were used.

Potential correlations between lung function parameters and ACT/CACT scores were tested using Pearson's r correlation for parametric variables and Spearman's rank for ordinal or non-parametric data. The above analyses were performed after converting spirometric parameters into z-scores using GLI software.

Paired t-tests and Wilcoxon signed rank tests were used to assess the change in ACT/CACT scores and number of UHAs in children at follow-up. Statistical analyses were performed using IBM SPSS Statistics for Windows (V.24.0, IBM, Armonk, New York, USA) and GraphPad Prism V7.00 for Windows (GraphPad Software, USA, www.graphpad.com). All statistical tests were performed at $\alpha=5\%$ level.

RESULTS

Participating practices

Ten practices participated in this study. They served a population of almost 120 000 people, ranging from ~3500 to 48 000 registered patients. Geographically, five of the practices were located in inner-city Leicester, three were in surrounding towns and two were in village locations.

Participants

Electronic database searches identified 1548 eligible children: 1097 (71%) were on their GPs asthma register and 451 (29%) were not, but had received asthma medications in the previous 12 months.

In total, 614 children responded to the written invitations and attended clinics held between June 2016 and August 2017; of these, 456 (75%) were on the asthma register. Sixty-three per cent of children were recruited in the months of March to August, and the remainder in the months of September to February.

Written informed consent was obtained from carers of 613 children. One parent refused consent and one parent later withdrew their consent without giving a reason, leaving 612 children in total. The characteristics of recruited children are shown in table 1. Overall, the participation rate was 40% of eligible patients, but ranged from 8% to 59% between practices. The practice recruiting only 8% of eligible children was an outlier, and due in part to recruitment at this practice commencing late into the study and therefore fewer clinics (in relation to the practice size) were set up compared with other practices.

Baseline spirometry, FeNO and asthma control test data

Spirometry

Acceptable spirometry was achieved by 575 (94.0%) children.

Using the definition FEV₁ or FEV₁/FVC < LLN, 135 out of 575 (23.5%) children had airflow obstruction. There was no difference in the prevalence of obstructed spirometry between children with diagnosed asthma versus those without a confirmed diagnosis, 23.7% vs 23.4% ($p=0.95$).

For comparison, the numbers of children defined as having airflow obstruction according to different thresholds are presented in table 2. Compared with using GLI LLN, using a fixed cut-off of FEV₁ < 80% predicted or FEV₁/FVC < 80% misidentified an additional 37 (6%) children as having airflow obstruction, but using the NICE recommended cut-off of FEV₁/FVC < 70% would have missed airflow obstruction in 108 (18%) children.

Table 1 Baseline characteristics of recruited children

	On asthma register (n=456)	Not on asthma register (n=156)	P value
No. of males (%)	247 (54.2)	85 (54.5)	0.945
Mean age in years (SEM)	10.3 (0.15)	9.1 (0.26)	<0.001*
No. of children with previous spirometry testing (%)	53 (11.7)	5 (3.2)	0.002*
Total daily prescribed dose (beclomethasone equivalent) of inhaled corticosteroids in micrograms (SEM)	227.41 (10.47)	87.18 (12.32)	<0.001*
Mean number of UHAs per child in preceding 6 months (SEM)	0.29 (0.03)	0.37 (0.06)	0.175
Number of children with ≥ 1 UHA in preceding 6 months (%)	98 (21.5)	42 (26.9)	0.163
Mean FEV ₁ % predicted (SEM)	93.29 (0.64)	93.44 (1.05)	0.904
Mean FEV ₁ z-score (SEM)	-0.45 (0.05)	-0.48 (0.09)	0.797
Mean FVC % predicted (SEM)	98.36 (0.69)	97.84 (1.00)	0.694
Mean FVC z-score (SEM)	-0.06 (0.05)	-0.13 (0.08)	0.462
Mean FEV ₁ /FVC (SEM)	83.75 (0.37)	84.29 (0.57)	0.451
FEV ₁ /FVC z-score (SEM)	-0.70 (0.05)	-0.66 (0.08)	0.701
Median FeNO in ppb (IQR)	25 (12–54)	16 (9–37)	0.004*

*Statistically significant difference ($p<0.05$) between subgroups. UHA, unplanned healthcare attendance.

Reversibility testing using salbutamol was performed in all children with FEV₁ < 80% predicted or FEV₁/FVC < 80% on baseline spirometry. Of the 172 children meeting this criterion, 56 (32.6%) demonstrated an increase in FEV₁ (L) of $\geq 12\%$ postbronchodilation.

Fraction of exhaled nitric oxide

FeNO results were obtained from 472 (77.1%) children. FeNO ≥ 35 ppb was identified in 171 children (36%) ranging from 36 to 231 ppb. The median (IQR) FeNO for children on the asthma register was higher; 25 (12–54) vs 16 (9–37), median difference = -5.0, 95% CI (-9.0 to 2.0), $p=0.004$.

Four hundred and sixty-five (76%) children were able to perform both spirometry and FeNO testing (figure 1). Only half had normal results for both tests, $n=240$ (52%).

Current asthma symptom control

Among our cohort, 256 children (41.8%) had poor current asthma symptom control according to their ACT/CACT test scores. One hundred ninety-seven out of 456 (43.2%) on the asthma register and 59 out of 156 (37.8%) not on their GP asthma register had ACT/CACT scores ≤ 19 ($p=0.24$).

Table 2 Number of abnormal spirometry results by definition used

Abnormality	Definition used	Number
Obstruction	FEV ₁ or FEV ₁ /FVC < LLN	135 (23.5%)
	FEV ₁ % predicted < 80 or FEV ₁ /FVC < 80%	172 (29.9%)
	FEV ₁ /FVC < 70%	27 (4.7%)
Restriction	Isolated FVC predicted < 80%	5 (0.9%)
	Isolated FVC < LLN	4 (0.7%)

LLN, lower limits of normal.

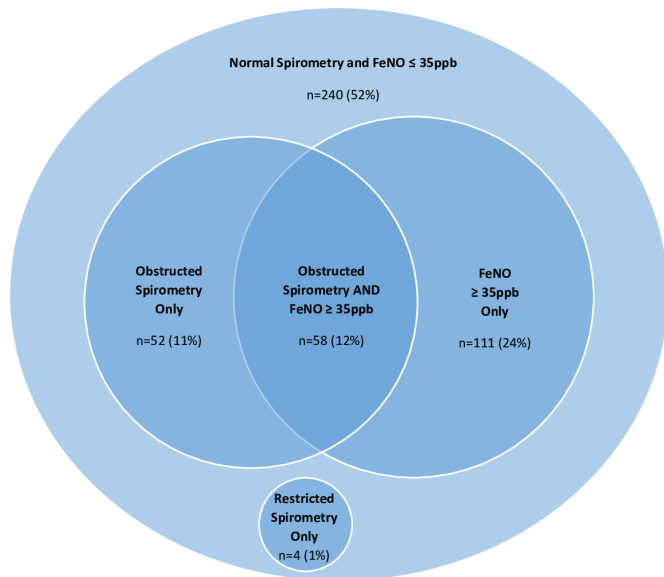


Figure 1 Number (percentage) of children with each combination of test results, based only on children who were able to perform both tests satisfactorily. Obstructed spirometry defined as FEV_1 or $FEV_1/FVC < LLN$ using GLI reference values. Restricted spirometry defined as $FVC < LLN$. GLI, global lung initiative; FeNO, fraction of exhaled nitric oxide; LLN, lower limits of normal; ppb, parts per billion.

Relationship between spirometry, FeNO and current asthma symptom control

Children with current uncontrolled asthma ($CACT/ACT \leq 19$) were more likely to have obstructed spirometry compared with those with good current control ($CACT/ACT > 19$); 28.5% vs 20.2% ($p=0.022$). Although both FEV_1 z-scores ($r=0.172$; 95% CI 0.037 to 0.302) and FEV_1/FVC z-scores ($r=0.238$; 95% CI 0.105 to 0.362) correlated weakly with ACT scores (children 12–16 years), the correlation coefficients were low, with a wide spread of data. In younger children (5–11 years), while there was also a weak trend towards higher values for FEV_1 and FEV_1/FVC with higher CACT scores, these did not reach significance (figure 2).

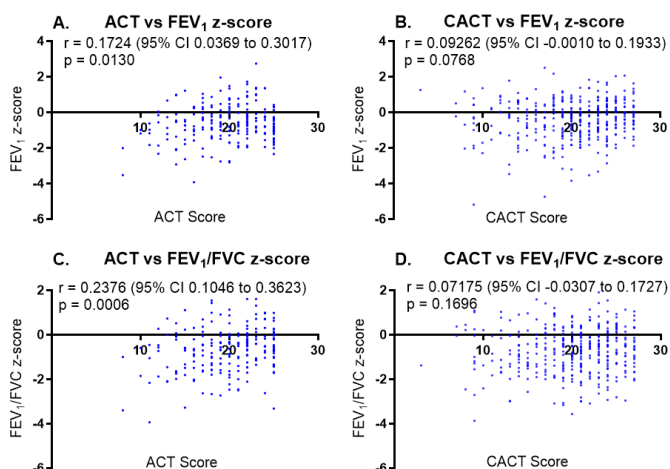


Figure 2 (A–D) Scatter plots showing the relationships between spirometric parameters and asthma control as measured using the asthma control test (ACT) for children 12 years and over, and the childhood asthma control test (CACT) for those aged 5–11 years. The correlation coefficient (r), CI values and p values are shown.

Table 3 Number of children with each combination of results according to reported current asthma symptom control

	Obstructed spirometry	Normal spirometry	Restricted spirometry
Poor control ($ACT/CACT \leq 19$), n=191			
FeNO ≥ 35	25 (13%)	46 (24%)	0
FeNO < 35	25 (13%)	93 (49%)	2 (1%)
Good control ($ACT/CACT > 19$), n=274			
FeNO ≥ 35	33 (12%)	65 (24%)	0
FeNO < 35	27 (10%)	147 (54%)	2 (1%)

Obstructed spirometry defined as FEV_1 or $FEV_1/FVC < LLN$ using GLI reference values. Normal spirometry defined as FEV_1 and $FEV_1/FVC \geq LLN$. Restricted spirometry defined as $FVC < LLN$.

ACT, asthma control test; CACT, childhood asthma control test; FeNO, fraction of exhaled nitric oxide.

There was no difference in the percentage of children with $FeNO \geq 35$ ppb between those with good or poor current asthma symptom control.

In children where both spirometry and FeNO data were available (table 3), 46% of children reporting good current control ($ACT/CACT$ score > 19) had at least one objective test abnormality. By contrast, in those reporting poor current control, 93 out of 191 (49%) children had normal spirometry and FeNO.

In terms of UHAs, children with poor current asthma control ($ACT/CACT \leq 19$) and obstructed lung function (GLI LLN) had almost double the mean (SEM) number of UHAs in the 6 months prior to review compared with children with poor current control but normal spirometry; 0.58 (0.11) vs 0.31 (0.05), mean difference = -0.27 , 95% CI (-0.07 to 0.47), $p=0.036$. There was no difference in the mean (SEM) number of UHAs in children with poor current control and raised FeNO versus children with poor current control and normal FeNO; 0.42 (0.09) vs 0.42 (0.07), mean difference = -0.001 , 95% CI (-0.23 to 0.23), $p=0.861$.

Potential role of spirometry and FeNO in asthma diagnosis

Spirometry and FeNO data were available from 109 children who had received asthma medications in the previous year, but who were not on the asthma register. Of these, 12% had obstructed spirometry (GLI LLN) and $BDR \geq 12\%$ to salbutamol and could have asthma confirmed according to NICE guidance. A further 6% had obstructed spirometry and $FeNO \geq 35$ ppb, 11% had obstructed spirometry only, and 13% had raised FeNO only; these children would warrant further investigation. Normal spirometry and FeNO was found in 58% of children not on the asthma register.

Follow-up

Unplanned healthcare attendances

Electronic records were reviewed for 605/612 children 6 months post asthma review. Seven children had moved GP surgery so their records were no longer accessible. The mean (SEM) number of UHAs fell from 0.31 (0.03) per child in the 6 months preceding review to 0.20 (0.02) per child over the 6 months following review; mean difference = -0.11 , 95% CI (-0.05 to 0.17), $p=0.0004$.

The number of children with at least one UHA also fell from 140 out of 612 (23%) children to 100 out of 605 (16.5%) children at follow-up, representing a relative reduction of 28%.

Current asthma symptom control

Follow-up asthma control tests were returned by 226 (37%) children. Overall, median (IQR) ACT scores improved from 20 (17–23) to 22 (19–24), median difference = -1.0 , 95% CI (-2.5 to 0.0), $p=0.032$, and CACT from 21 (19–24) to 23 (19.5–25), median difference = -1.5 , 95% CI (-2.0 to 0.5), $p<0.0001$ in the 6 months following review.

DISCUSSION

We conducted a large study in primary care to explore the feasibility of spirometry and FeNO testing for the management of children's asthma. Abnormal spirometry was identified in almost one-quarter of children studied. Although comparison of prevalence of abnormal spirometry with other studies is not straightforward, due to different criteria for normality and reference equations used, our finding is broadly similar to two North American studies involving children with asthma.^{15 16} Previous studies have shown that children with an FEV₁ of 60%–80% predicted are 1.4–1.8 times more likely to experience an asthma attack over the subsequent year.^{17 18}

FeNO levels ≥ 35 ppb were identified in 36% of our cohort. This has potential implications for predicting the risk of future loss of asthma control,¹⁹ and FeNO-guided asthma management may have a role in reducing asthma attacks.⁶

Given the association between abnormal spirometry and FeNO with poor asthma outcomes, the high prevalence of abnormal lung function and FeNO seen in our cohort highlights the need for objective measurements in asthma monitoring to fully appreciate the future risk of adverse asthma events.

In children able to perform both spirometry and FeNO testing, around half had at least one abnormal test result. Of those with at least one abnormality, approximately half had raised FeNO alone, one-quarter had obstructed spirometry alone and one-quarter had abnormal results for both, suggesting a complementary role for both FeNO and spirometry in children's asthma monitoring.

We found a statistically significant, but weak relationship between ACT scores with FEV₁ and FEV₁/FVC, and no correlation between CACT scores with either FEV₁ or FEV₁/FVC. There was also no correlation between FeNO with either ACT or CACT scores. Recent studies have similarly reported weak correlations between spirometry and ACT in adults,²⁰ but not between spirometry or FeNO and CACT in younger children.²¹ This might reflect a disassociation between a patient's asthma symptoms and their perception of those symptoms,^{22 23} or a temporal delay between changes in symptoms and lung function.^{24 25}

Importantly, almost half the children in our cohort reporting good current asthma control had at least one objective test abnormality; with 12% having both abnormal spirometry and FeNO. This suggests that asthma severity²⁶ and suboptimal control² may be under-recognised when clinical evaluation is based on symptoms alone. By contrast, 49% of children reporting poor current control had normal tests, suggesting either incorrect asthma diagnosis, or highlighting that children can be symptomatic despite normal lung function. Similarly, a US study involving 201 children found that asthma assessments based on history alone underestimated asthma severity determined by spirometry in 31% of children, while assessments using spirometry alone under-recognised poor control in 40% of symptomatic children compared with clinician-determined severity.²⁷ Although this study predates publication of GLI reference values, making direct comparison with our data challenging, it highlights the

fact that asthma assessments based on either symptoms, or objective tests in isolation do not provide a full picture of the child with asthma.

Under-recognition of the underlying severity of a child's asthma has potentially severe consequences. Two recent coroners' reports in the UK, both identified failures in the management of two children who died in 2014 and 2015, emphasising the lack of appreciation and recognition of the severity of their condition and the deteriorating nature of their clinical control (Carney 2015; Radcliffe 2017). We believe this highlights the need for objective measures of asthma control in addition to the traditional symptoms-based approach.

Around a quarter of our cohort had at least one UHA in the 6 months preceding review. This is higher than previously reported asthma attack rates in children of 8%–36% per year^{28 29} and may be due to different definitions for asthma attacks. Previous UK studies^{28 30} defined attacks as ED attendances or the need for oral corticosteroids (OCS); these criteria may have missed milder attacks where no OCS were prescribed. We included any asthma-related UHA regardless of OCS prescription where children received increased inhaled or nebulised short-acting bronchodilators, and referred to discharge letters stored as text documents even if they were not coded electronically. The recent *Lancet* commission on asthma³¹ discusses the heterogeneity of asthma attacks at all ages and emphasises that any loss of symptom control should be taken seriously.

In children reporting poor current asthma control, we found those with obstructed airflow had almost double the mean number of UHAs at baseline compared with children with normal lung function. This association has potentially important implications for asthma attack risk stratification.

Notably, following an asthma review that included objective tests, we observed a small improvement in ACT/CACT scores during follow-up. This is arguably subjective and possibly a result of study (Hawthorne) effect.³² Moreover, the improvement in asthma control test scores at follow-up were below the published minimally clinically important change in scores for both ACT and CACT of 3-points and 2-points, respectively.

However, the mean number of UHAs and the proportion of children having at least one UHA also fell by almost a third. We speculate that this resulted from practice nurses being trained as part of this study to act on both patient-reported symptoms and objective evidence from spirometry and FeNO, leading to an increased recognition of suboptimal control, undertreatment or poor compliance with preventer medications and more accurate titration of asthma treatment. This requires confirmation in an adequately powered clinical trial. Previous studies have similarly demonstrated that asthma reviews conducted by trained asthma nurses in primary care can improve patient outcomes and reduce asthma attacks^{33 34}; however, these studies did not include the use of spirometry or FeNO as part of the asthma review.

Previous asthma studies comparing standard symptoms-based monitoring versus monitoring with additional spirometry or FeNO have not found significant differences between groups in terms of reported symptoms. There is, however, some evidence that children monitored with FeNO may have fewer asthma attacks.^{6 35 36}

An Australian trial comparing standard asthma monitoring versus active monitoring with spirometry reported no difference in number of asthma attacks.³⁷ This trial did not however, include a management algorithm to direct treatment of children with abnormal lung function. Intuitively, spirometry-based monitoring would only be effective if abnormal results are acted on appropriately.

In terms of diagnosis, one-quarter of our cohort were identified based on them receiving asthma medications in the previous year despite not having a coded asthma diagnosis. We believe this may reflect both a hesitance to formally diagnose children with asthma in primary care, and an issue with coding practices, that is, the child is diagnosed but this is not recorded on the practice database. These children would not automatically be recalled for a routine annual asthma review and are at risk of having unrecognised poor symptom control. Using the NICE asthma diagnosis algorithm, 12% of our cohort without an existing asthma diagnosis had asthma confirmed, and a further 6% had probable asthma based on two positive tests, demonstrating the potential usefulness of providing objective tests within primary care. This is similar to the findings of the NICE asthma guideline feasibility study (NICE 2017), which were able to confirm an asthma diagnosis in 24.5% of their cohort, who presented acutely with asthma symptoms. As both lung function and FeNO can fluctuate, it is possible that our diagnosis rate was slightly lower, due to our cohort of children being recruited during routine reviews and not when they were presenting acutely unwell.

Strengths: we prospectively recruited a large cohort of children from practices representative of different ethno-socioeconomic populations. All but two children attending the asthma review participated. As follow-up data were collected primarily from electronic records, only seven (1%) children were lost to follow-up; in remaining children, datasets were complete for baseline and follow-up UHAs, making our data robust. All training for spirometry and FeNO were delivered by the same study team members; and every spirogram was reviewed for acceptability following ATS/ERS recommendations to ensure quality of lung function data.

Limitations: the absence of a control arm makes the interpretation of the apparent fall in asthma attacks seen in our cohort difficult. As attendance was voluntary, it is possible that only more symptomatic children attended; biasing our cohort towards those with poorer current asthma symptom control and poorer lung function. Second, practice staff were trained to interpret spirometry using fixed cut-offs rather than GLI LLN, despite the latter being the gold standard. This was a conscious decision by the study team for pragmatic reasons, as practice staff were already familiar with the use of fixed cut-offs in the context of performing adult spirometry. As routine spirometry becomes accepted in general practice, we recommend that fixed cut-offs should be superseded by LLN.

Also, asthma is a chronic relapsing condition and this study has included clinical assessments at one point in time only; therefore, our findings and conclusions relate to that one snapshot and may not accurately reflect longitudinal asthma control. Additionally, we did not set out to control for the effect of seasonality on symptom control and asthma attacks, however we recruited our patients over a 14-month period across all four seasons, making a seasonal influence on our data unlikely.

Finally, all participating practices were from the same, although diverse, geographical region within the East Midlands, UK; it is possible that only the most motivated practices expressed an interest to participate, resulting in a group of practices which may not be representative.

To our knowledge, this is the first primary care study in the UK to explore the role of spirometry and FeNO testing in monitoring children with asthma. We have shown that abnormal lung function and FeNO are highly prevalent in children who attend for asthma reviews in primary care, and correlates poorly with patient-reported symptom scores. Symptomatic children with abnormal lung function were also more likely to have had an

asthma attack during the study period. Moreover, we observed a reduction in the number of asthma attacks and improved current asthma control test scores in the 6 months after the clinical review which included objective tests. This important observation requires confirmation with an adequately powered randomised clinical trial, with repeat objective testing over time, in order to demonstrate causation and longitudinal changes in lung function. To conclude, our data suggests that a symptoms-based assessment alone is inadequate, and is likely to miss children at high risk of a near future severe asthma attack.

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