

Role of inducible nitric oxide synthase in asthma risk and lung function growth during adolescence

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ABSTRACT

Background Inducible nitric oxide (NO) synthase (iNOS, encoded by *NOS2A*) produces NO in response to environmental stimuli, which can result in nitrosative stress. Because nitrosative stress affects respiratory health, it was hypothesised that variants in *NOS2A* are associated with asthma incidence and lung function growth during adolescence.

Methods In this prospective study, spirometric testing was performed at school and a presence or absence of asthma was ascertained annually by questionnaire among children participating in the Southern California Children's Health Study. 24 single nucleotide polymorphisms (SNPs) of the *NOS2A* region (with seven promoter SNPs in one haplotype block), spanning 20 kb upstream and 10 kb downstream were genotyped. Association between the *NOS2A* region and asthma or lung function growth was tested using genetic block-specific principal component and haplotype analyses. This study was restricted to children with Latino and Caucasian ancestry for analyses of both asthma (n=1596) and lung function growth (n=2108).

Result A pair of "yin-yang" haplotypes in the promoter region showed strong association with new-onset asthma and lung function growth. The "yin" haplotype (h0111101) was associated with 44% increased asthma risk (p=0.003) and reduced forced expiratory volume in 1 s (FEV₁) growth from 10 to 18 years of age (-29.46 ml, p=0.07), whereas the "yang" (h1000010) haplotype was associated with 23% reduced asthma risk (p=0.13) and better FEV₁ growth (43.84 ml, p=0.01). Furthermore, the increased asthma risk associated with h0111101 was restricted to children with the *GSTM1* "null" genotype (interaction p=0.002, HR 1.89, 95% CI 1.34 to 2.60).

Conclusion Common haplotypes in the *NOS2A* promoter are associated with new-onset asthma and lung function growth. These effects are stronger in adolescents with the *GSTM1* "null" genotype.

INTRODUCTION

Studies have shown that oxidative stress is an important determinant of respiratory health.¹⁻³ Both increased incidence of asthma and reduced lung function growth during adolescence have been associated with environmental and genetic determinants of oxidative stress.^{4,5} Like reactive oxygen species (ROS), reactive nitrogen species (RNS) can also produce cellular injury through nitrosative stress.⁶ Inducible nitric oxide synthase (iNOS) plays an important role in determining nitrosative stress, as it produces large amounts of nitric oxide (NO) in response to environmental stimuli.⁶

NO is a gaseous signalling molecule that is involved in a spectrum of biological processes. In response to proinflammatory cytokines and lipopolysaccharides, epithelial cells and T lymphocytes produce NO through iNOS.⁶ NO reacts with superoxides (O₂·) to form the highly reactive peroxynitrites (ONOO⁻) that have been shown to produce airway inflammation⁷ and cause airway remodelling.^{8,9} Furthermore, the fractional concentration of NO in exhaled breath, a marker of airway inflammation,¹⁰ is predominantly of iNOS origin.¹¹ Increased expression of iNOS may play a role in inflammation of the upper and lower airways, and contribute to both allergic rhinitis and asthma.¹²

Despite the evidence supporting the potential importance of iNOS in respiratory health, limited information is available about the role of DNA sequence variations in *NOS2A* that encodes iNOS. *NOS2A* is located at chromosome 17q11.2-q12, a region identified by genome-wide association tests as a possible region determining asthma and atopy.¹³ A large number of variants in *NOS2A* have been identified that have the potential to affect expression or function. The induction of *NOS2A* is complex, and the promoter region, which contains a number of variants, extends up to 16 kb upstream of the gene.¹⁴ There is limited evidence that repeat polymorphisms in the promoter region are associated with atopic conditions.¹⁵⁻¹⁸ In addition to the repeat polymorphisms, single nucleotide polymorphisms (SNPs) also exist in the promoter region that are not in strong linkage disequilibrium (LD) with the repeat polymorphisms.¹⁹ To date, a single study involving Czech adults reported association between SNPs in the *NOS2A* promoter region and atopy and asthma severity.¹⁶

In this study, we investigated the role of variations in the *NOS2A* gene with asthma pathogenesis and lung function growth in children. We hypothesised that genetic variants of *NOS2A*, especially those in the promoter region, are associated with new-onset asthma during adolescence and lower lung function growth, due to their likely effect on nitrosative stress. Assessing the effects of the *NOS2A* variants across these two respiratory outcomes provided an opportunity to evaluate the consistency in the findings. Furthermore, because the formation of peroxynitrite is dependent on the availability of ROS, we hypothesised that the associations between *NOS2A* polymorphisms and respiratory outcomes vary by genetic determinants of oxidative stress (*GSTM1*, *GSTP1*, *CAT* and *HMOX1*) that previously have been reported to be

associated with lung function growth or asthma pathogenesis in this cohort.^{4 20 21} We investigated these hypotheses in a cohort of Latino and Caucasian children who participated in the Children's Health Study (CHS).^{22 23} Some of the results have been reported in abstract form.²⁴

METHODS

Further details are available in the online supplement.

Subjects and materials

Details concerning the design, methods and characteristics of the CHS cohort have been presented previously.^{22 23} Briefly, in 1993, fourth-, seventh- and tenth-grade children were enrolled into the study in each of 12 communities in Southern California. A second cohort of fourth-grade children was recruited from the same schools in 1996. Study subjects were followed annually until high school graduation. Detailed health and socio-demographic data were collected annually.

As in previous CHS genetic analyses,⁴ this study was restricted to children of Latino and Caucasian ancestry with available genetic data. The analysis of new-onset asthma included 1596 children who were free of asthma and wheeze at baseline. Similarly,²⁵ the analysis of lung function was restricted to the two cohorts of fourth-grade children (mean age 10 years (SD=0.44)) who were recruited in 1993 (cohort 1, n=912) and 1996 (cohort 2, n=1196) and had similar lung function growth patterns.

New-onset asthma

Children with no prior history of asthma at study entry who subsequently reported physician-diagnosed asthma at annual follow-up were classified as having new-onset asthma. Children were also interviewed annually about the use of inhaled medications. We defined a restricted group of new-onset cases with recent inhaler use for sensitivity analyses (n=116).

Lung function measurement

Data on children's lung function were collected by trained field technicians using a standardised protocol during annual school visits. Maximal-effort spirometry and standing height and weight were measured. Details regarding the lung function testing protocol have been published previously.²² Three measures of lung function were included in this analysis: forced vital capacity (FVC), forced expiratory volume in the first second (FEV₁) and forced expiratory flow over the mid-range of expiration (FEF₂₅₋₇₅).

NOS2A haplotype block determination

Most of the upstream regulatory and promoter region was contained in a single block (figure 1: block 7) with a low multiallelic D' (0.34) with the adjacent block (figure 1: block 6). Blocks 1–6 had higher multiallelic D' (>0.70). Therefore, the remaining region (blocks 1–6) was analysed as two segments. The first segment was comprised of blocks 1–3 and the second segment of blocks 4–6 (figure 1).

Haplotype frequencies of unphased NOS2A SNPs of the promoter region and the two coding segments for Latino and Caucasian subjects were estimated using tagSNPs. The estimated number of copies of each haplotype was used as a proxy for the true haplotype, a single imputation procedure that provides unbiased estimates and appropriate CIs. Four haplotypes in block 7 explained at least 97% of the variability of this block in both ethnic groups. Furthermore, a "yin–yang" pair of haplotypes (h0111101 and h1000010), two high-frequency haplotypes that

have completely mismatching SNP nucleotides at every SNP location,²³ explained at least 65% of the variability of this promoter block in the Latino and Caucasian population.

Statistical methods

To assess the global association of variation across the locus with asthma occurrence or lung function growth, the groups of common SNPs and haplotypes were tested using both principal component (PC)³¹ and haplotype approaches. The statistical significance was based on the likelihood ratio tests (LRTs) comparing the full model (model with genetic data adjusted for all necessary adjustment variables) with the base model (model with all necessary adjustment variables without any genetic data). As indicated by Gauderman *et al*, a block-specific PC analysis was preferred over a whole locus approach.³¹ PCs explaining at least 80% of the variance of each of the three regions (two PCs for segment 1, four PCs for segment 2 and one PC for the promoter region block (block 7)) were modelled to determine the overall statistical significance of each region across outcomes (figure 1).

The haplotype model consisted of all the haplotypes with frequency >5% and "other" haplotype, with the most common haplotype as the reference group. We also tested the effect of one and two copies of the identified haplotypes across the respiratory outcomes, using all other haplotypes as the reference group. After identifying a region that showed an overall statistically significant association with the respiratory outcomes, we tested associations between individual SNPs contained in that region and the respiratory outcomes. To address the issue of multiple testing of the correlated SNPs, the single SNP analyses were adjusted for multiple testing using the p_ACT method.³²

New-onset asthma analysis

We fitted Cox proportional hazard regression models, with the time scale defined as the follow-up time with sex- and age- (integer years) specific baseline hazards. All models were adjusted for community and ethnicity markers. Associations are expressed as HR and 95% CIs.

Lung function analysis

A hierarchical mixed effects model was used to relate 8-year growth in each lung function measure to NOS2A, with a basic structure that has been previously described.³ Random effects for the intercept and 8-year growth parameters were included at the subject level. We estimated and tested the effect of NOS2A on 8-year lung function growth from age 10 to 18. Because the findings were similar for all three lung function measures (online table E5), we present the results for FEV₁ only.

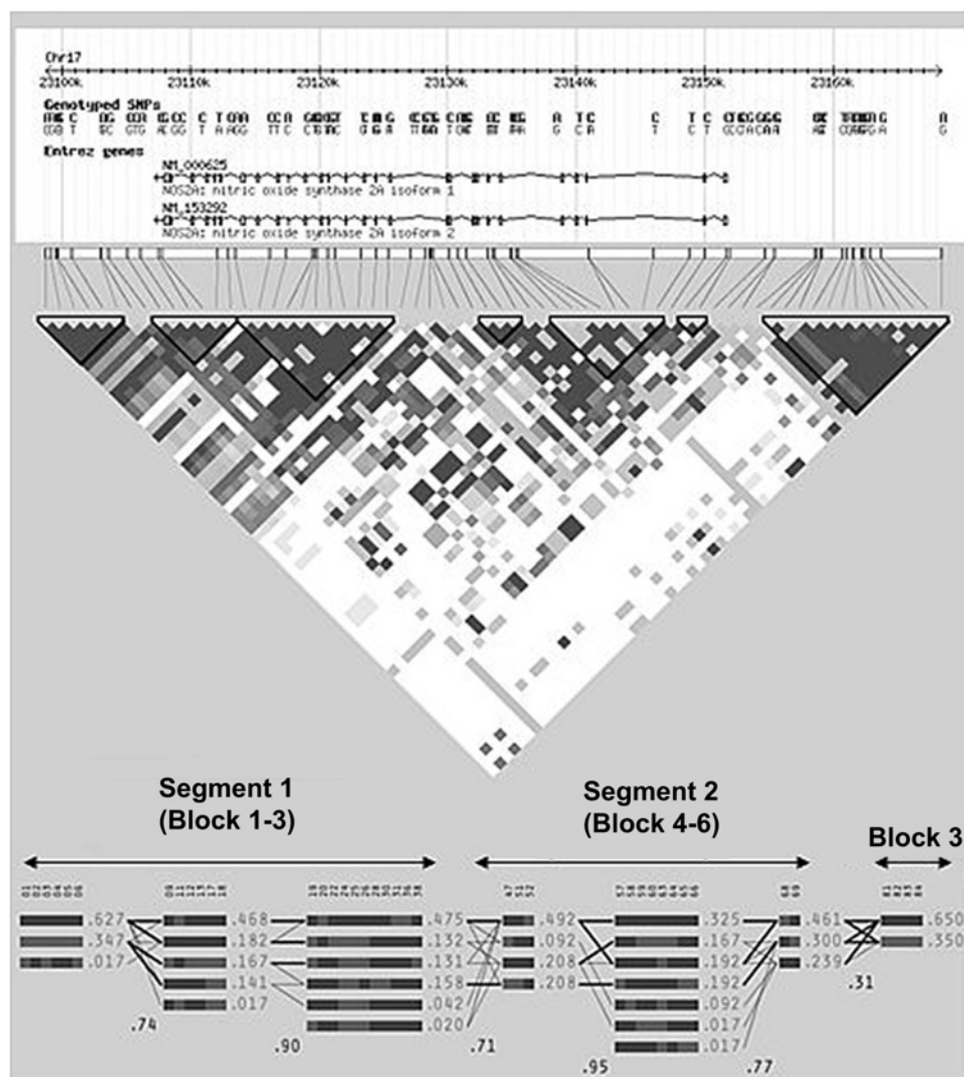
Sensitivity analyses

Other variables available from questionnaire data were evaluated for potential confounding in both asthma and lung function models, but ultimately removed from the final analysis since they did not alter the results. Based on our previous publications, we also considered the "null" deletion of GSTM1,²⁰ GSTP1-Ile105Val,²⁰ (GT)_n repeats of HMOX1 and CAT-262C→T⁴ as possible confounders.

Test of heterogeneity

To test whether other genes involved in the oxidative pathway (GSTM1, GSTP1:Ile105Val, HMOX1 and CAT) modified the association between NOS2A and respiratory outcomes, models with and without appropriate interaction terms were considered. In the presence of statistically significant heterogeneity among subgroups, stratified analysis was performed.

Figure 1 The linkage disequilibrium (LD) plot and haplotypes of the *NOS2A* gene (including the promoter region) for Europeans using data from the International HapMap Project. For the LD plot, the triangles represent the haplotype blocks. Haplotype frequencies within each block are shown below the blocks, and the multiallelic D' between blocks are shown between haplotypes.



All hypothesis testing was conducted assuming a 0.05 significance level and a two-sided alternative hypothesis. All analyses were conducted using SAS software Version 9.1.3 (SAS Institute, Cary, North Carolina, USA). We used `p_ACT_seq` R program in R software (V 2.21) to address the issue of multiple testing (http://csg.sph.umich.edu/boehne/p_act.php).

RESULTS

Baseline characteristics

The majority of children included in the asthma ($n=1596$) or lung function ($n=2108$) analyses were 9 years of age at study entry and of Caucasian descent (table 1). Compared with children in the asthma analysis, those in the lung function analysis were more likely to have family history of asthma (21.7% vs 13.9%) and history of atopy (48.4% vs 35.5%) at baseline as the asthma analysis was restricted to children without any history of wheeze or asthma diagnosis at baseline. We did not observe any other significant differences between the two populations at baseline, and 998 children were common to both analyses.

One hundred and fifty cases of new-onset asthma were diagnosed during the follow-up period. The crude incidence rate (IR) of asthma was 16.3/100 person-years and showed little difference between Caucasians (IR=16.1/1000 person-years) and Latinos (IR=16.6/1000 person-years). The prevalence of wheeze

and prevalent asthma in the lung function analysis group was 34.7% and 15.0% at baseline, respectively. The mean FEV₁ growth for boys and girls over the 8-year follow-up time was 2441 ml and 1367 ml, respectively.

MAIN EFFECT OF *NOS2A*

Utilising a PC-based analysis for global tests, we observed that block 7 (containing seven SNPs of the promoter region) was associated with both an increased risk of new-onset asthma ($p=0.002$) and decreased FEV₁ growth ($p=0.02$) during adolescence (online table E6). The haplotype-based analysis for global association also showed a similar pattern of association (table 2). Compared with the common haplotype, h0111101, all other haplotypes were protective for new-onset asthma. The h0111101 haplotype was associated with increased risk of new-onset asthma (HR 1.44, 95% CI, $p=0.003$) and lower FEV₁ growth (by 29.5 ml, 95% CI, $p=0.07$) compared with all other haplotypes (table 3). This haplotype also showed a dose-dependent effect for asthma risk ($p=0.01$). Children with one or two copies of this haplotype were at 1.49-fold (95% CI 1.03 to 2.14) and 2.08-fold (95% CI 1.25 to 3.45) increased risk of new-onset asthma, respectively, compared with children without this haplotype. The other member of the “yin–yang” pair, h1000010, was associated with higher 8-year FEV₁ growth ($p=0.01$) and showed a dose-dependent pattern ($p=0.02$).

Table 1 Selected baseline characteristics of children who were included in the new-onset asthma and lung function analyses

	Children included in incident asthma analysis (n=1596)	Children included in LF analysis (n=2108)
Age group		
7–9 years	875 (54.8)	1603 (76.0)
10–11 years	335 (21.0)	505 (24.0)
>11 years	386 (24.2)	–
Sex		
Girls	894 (56.0)	1086 (51.5)
Boys	702 (44.0)	1022 (48.5)
Latino	554 (34.7)	710 (33.7)
Overweight	395 (24.7)	536 (26.5)
Family history of asthma	222 (13.9)	430 (21.7)
History of atopy*	567 (35.5)	916 (48.4)
History of asthma at study entry	0 (0)	310 (12.0)
Household ETS exposure	186 (19.0)	361 (17.9)
In utero exposure to smoking	219 (13.7)	354 (17.2)
Pests of any kind	1195 (74.9)	1627 (83.4)
Pets of any kind	1302 (81.6)	1728 (82.0)
Children with health insurance	1313 (82.3)	1781 (86.3)
Annual family income (in US dollars)†		
≤14 999	174 (10.9)	263 (14.3)
15 000–49 999	584 (36.6)	768 (41.7)
≥50 000	597 (37.4)	812 (44.1)
Highest parental education level‡		
Less than high school	210 (13.2)	222 (10.9)
College	1122 (70.3)	1567 (74.3)
Graduate	208 (13.0)	242 (11.9)

*Children with any history of allergy, hay fever or rhinoconjunctivitis were defined as “atopic”.

†Numbers do not add up to 100% due to missing information. ETS, environmental tobacco smoke; LF, lung function.

Compared with children not carrying this haplotype, those with one or two copies of this haplotype had 28.6 ml (95% CI –15.62 to 72.93) or 107.7 ml (95% CI 32.54 to 183.93) higher

FEV₁ growth over an 8-year period. A marginally significant protective trend for new-onset asthma was also observed among carriers of this haplotype. The risk of asthma decreased by 14% and 54% among carriers of one or two copies of the h1000010 haplotype.

SNP ANALYSIS

Investigation of the single SNPs corresponding to the haplotypes for the upstream promoter region (block 7) demonstrated significant associations (online table E7) consistent with haplotype results; therefore, we report the results from h0111101 and h1000010 analyses only.

Sensitivity analysis

The observed association between h0111101 and respiratory outcomes (new-onset asthma and FEV₁ growth) were not substantially affected by adjustment for potential confounders (online tables E8 and E9: model 1). The associations between h0111101/h1000010 and respiratory outcomes were not confounded by the variants of genes involved in oxidative stress and asthma—that is, *GSTM1*, *GSTP1*, *HMOX1* and *CAT*⁴ (online tables E8 and E9: model 2).

To assess whether the effect of the haplotypes on FEV₁ growth was due to reduced lung function level in children with asthma, we investigated the effects only among children without prevalent or new-onset asthma (n=1672). In this asthma-free population, the beneficial effect of h1000010 remained mostly unchanged (49.3 ml increase in FEV₁ growth; 95% CI 10.9 to 87.7); however, the detrimental effect of h0111101 was less pronounced (15.3 ml decrease in FEV₁; 95% CI –53.9 to 23.2).

To investigate whether the associations were consistent in independent groups of children, we performed stratified analyses for the two fourth-grade cohorts in the study populations independently recruited in 1993 and 1996. These cohorts were from the same communities and schools and thus had similar environmental exposure and socio-economic characteristics. The effect estimates for both new-onset asthma and FEV₁ growth

Table 2 Overall associations between *NOS2A* and new-onset asthma or FEV₁ growth (in ml) assessed using haplotype-based analysis

<i>NOS2A</i> region	Haplotype	New-onset asthma	8-year growth in FEV ₁		
		HR (95% CI)†	p Value§	Estimate (95% CI)‡	p Value§
Segment 1 (Blocks 1–3)	h0110010	Ref.	0.06	Ref.	0.46
	h1000000	0.72 (0.51 to 1.01)		9.57 (–31.99 to 51.14)	
	h1001101	1.30 (0.93 to 1.8)		27.64 (–19.76 to 75.04)	
	h0000000	1.00 (0.71 to 1.42)		12.40 (–34.78 to 59.57)	
	Other*	0.92 (0.62 to 1.37)		–25.24 (–75.13 to 24.65)	
Segment 2 (Blocks 4–6)	h0101100010	Ref.	0.25	Ref.	0.71
	h1010111001	1.21 (0.81 to 1.81)		–6.64 (–61.13 to 47.84)	
	h0010000100	0.66 (0.41 to 1.05)		31.86 (–21.68 to 85.40)	
	h0010000101	1.30 (0.84 to 2.03)		–12.89 (–74.98 to 49.20)	
	h0101000010	0.79 (0.48 to 1.30)		–32.94 (–92.81 to 26.92)	
	h0000000100	1.13 (0.73 to 1.74)		–2.89 (–61.28 to 55.51)	
	h1001011001	1.07 (0.63 to 1.82)		31.14 (–41.84 to 104.12)	
	h0001000001	0.99 (0.49 to 1.99)		10.34 (–82.78 to 103.47)	
	Other*	0.96 (0.64 to 1.45)		1.07 (–55.16 to 57.30)	
Block 7	h0111101	Ref.	0.02	Ref.	0.10
	h1000010	0.66 (0.49 to 0.88)		48.90 (11.60 to 86.20)	
	h0000010	0.84 (0.60 to 1.18)		3.86 (–41.95 to 49.68)	
	h0000000	0.62 (0.44 to 0.88)		17.04 (–26.77 to 60.85)	
	Other*	0.85 (0.42, 1.74)		1.30 (–107.56 to 110.17)	

*Haplotypes with <5% frequencies are grouped into the “Other” category.

†HR and 95% CI are based on fitting the Cox proportional hazard model (for details see the Methods section).

‡Estimate and 95% CI are based on hierarchical mixed effects model (for details see the Methods section).

§p Values are based on a likelihood ratio test comparing models with and without genetic data.

FEV₁, forced expiratory volume in 1 s.

were mostly similar across cohorts (online tables E3 and E4). In each of the cohorts and ethnic groups, h0111101 (online table E3) was associated with increased risk of new-onset asthma (HR range 1.2–1.7) and lower FEV₁ growth (FEV₁ growth range –13.8 to –51.8 ml). In contrast, h1000010 (online table E4) was associated with decreased risk of new-onset asthma (HR range 0.68–0.89) and higher FEV₁ growth in all the cohorts and ethnic groups (FEV₁ growth range 32.4–54.3 ml).

Joint effects with *GSTM1* “null” status

Lastly, we evaluated whether the association between the haplotypes of interest and respiratory disease was modified by *GSTM1* status. The association between the h0111101 haplotype and new-onset asthma varied by *GSTM1* “null” genotype ($p = 0.002$, online table 4). The increased risk of new-onset asthma for h0111101 carriers was restricted to children who lacked *GSTM1* (HR 1.89, 95% CI 1.3 to 2.6). For FEV₁, those with the h0111101 haplotype and *GSTM1* “null” genotype were also associated with the worst growth pattern (FEV₁ growth –53.19, 95% CI –99.63 to –6.75). The protective effect of h1000010 on asthma risk or FEV₁ growth did not vary by *GSTM1* genotype. The haplotype effect was not modified by the presence of functional variants in other oxidative stress genes—that is, *GSTP1*, *HMOX1* or *CAT*.

DISCUSSION

In this prospective study of new-onset asthma and lung function growth during childhood, we found that DNA sequence variation in the promoter region of *NOS2A* plays a potentially important role in respiratory health and development. We identified a pair of “yin–yang” haplotypes (h0111101 and h1000010) that accounted for >65% of the variation in the promoter region. The “yin” haplotype (h0111101) was associated with increased risk for asthma and poor lung function growth. In contrast, the “yang” haplotype (h1000010) was associated with higher lung function and lower asthma risk. Furthermore, our analyses suggest that promoter variation in *NOS2A* has independent effects on asthma occurrence and lung function. These effects were consistent across two different respiratory outcomes, different internal cohorts and two different ethnic groups.

Although *NOS2A* has been of interest in asthma research, most of the studies did not use SNPs and none of the studies investigated its role on lung function growth. Most of the studies involved a putative functional repeat polymorphism of the promoter region, (CCTTT)_n, that was associated with atopic status in Japanese adults,¹⁷ but not in Chinese children,¹⁸ Czech

adults¹⁶ or in Indian children and adults.¹⁵ In the present study, we found no evidence for an effect of this repeat polymorphism on asthma or lung function growth (data not shown). To the best of our knowledge, only one previous study has examined the role of SNPs in the promoter region on respiratory health, and associations with atopy and asthma severity were reported.¹⁶ Thus it is hard to compare our current finding with previous publications on *NOS2A*. However, the consistent effects of the “yin–yang” haplotypes for the two different respiratory outcomes suggest that more than one variant in this region may underlie the observed associations with asthma and lung function.

Besides being consistent across outcomes, the effects of the haplotypes were also independent of the respiratory outcomes. The detrimental effect of h0111101 and beneficial effect of h1000010 on lung function were observed among children with and without a history of asthma. Although the magnitude of the effect of the haplotypes was associated with an apparently small effect (1% change in lung function growth during adolescence in boys and 2% in girls), they may nevertheless play an important role in susceptible populations with low lung function levels. Furthermore, given the multifactorial nature of lung function growth and asthma, any genetic or environmental risk factors are not expected to have large individual effects.

Accumulating evidence indicates that nitrosative stress, like oxidative stress, plays an important role in airway pathobiology.⁷ Because the effects of variants in *NOS2A* on respiratory health may depend on levels of oxidative stress and gene variants in oxidant defence pathways modulate levels of oxidative stress,³³ we hypothesised that *NOS2A* may interact with variants in oxidant defence genes such as *GSTM1*, which have also been associated with asthma occurrence²¹ and lung function growth.²⁰ Our observation that the detrimental effect of h0111101 on asthma risk was most apparent among children with a common *GSTM1* variant lacking enzyme activity supports this hypothesis. If h0111101 results in increased NO production and formation of peroxynitrites, as we have speculated, a larger effect might be expected in *GSTM1* null individuals. Lower NO production in h1000010 might result in a protective effect irrespective of *GSTM1* status. Although levels of oxidative stress may be important, other pathways that involve *GSTM1* may also contribute. In the current study, the *NOS2A* associations were independent of variation in *GSTP1*, *HMOX1* and *CAT*, genes that are also involved in the oxidant defence pathways and have been shown to be associated with new-onset asthma and lung function. This implies that other functions of *GSTM1*, such as electrophile conjugation with

Table 3 Association between *NOS2A* promoter “yin–yang” haplotypes and new-onset asthma or FEV₁ growth (in ml)

<i>NOS2A</i> region	Haplotype	New-onset asthma		8-year growth in FEV ₁	
		HR (95% CI)*	p-Value‡	Estimate (95% CI)†	p Value‡
h0111101	None	Ref.	0.003	Ref.	0.07
	At least one	1.44 (1.13 to 1.82)		–29.46 (–61.27 to 2.36)	
h0111101	None	Ref.	0.01	Ref.	0.15
	One copy	1.49 (1.03 to 2.14)		–40.69 (–85.04 to 3.66)	
	two copies	2.08 (1.25 to 3.45)		–46.82 (–118.29 to 24.65)	
	None	Ref.	0.06	Ref.	0.01
h1000010	At least one	0.77 (0.59 to 1.01)		43.84 (11.39 to 76.28)	
h1000010	None	Ref.	0.09	Ref.	0.02
	One copy	0.86 (0.61 to 1.21)		28.65 (–15.62 to 72.93)	
	Two copies	0.46 (0.21 to 1.00)		107.74 (31.54 to 183.93)	

*HR and 95% CI are based on fitting the Cox proportional hazard model (for details see the Methods section).

†Estimate and 95% CI are based on the hierarchical mixed-effects model (for details see the Methods section).

‡p Value is based on a likelihood ratio test comparing models with and without genetic data.

FEV₁, forced expiratory volume in 1 s.

Table 4 Association between the *NOS2A* promoter haplotypes and new-onset asthma or FEV₁ growth, by *GSTM1* genotype

Haplotype	<i>GSTM1</i>	New-onset asthma		8-year growth in FEV ₁		
		HR (95% CI)*	Interaction p value‡	Estimate (95% CI)†	Interaction p value‡	
h0111101	None	Present	Ref.	0.002	Ref.	0.32
	At least one	Present	0.89 (0.58 to 1.37)		−4.64 (−50.20 to 40.91)	
	None	Null	Ref.		Ref.	
	At least one	Null	1.89 (1.34 to 2.60)		−53.19 (−99.63 to −6.75)	
h1000010	None	Present	Ref.	0.42	Ref.	0.53
	At least one	Present	0.85 (0.54 to 1.34)		39.85 (−5.81 to 85.50)	
	None	Null	Ref.		Ref.	
	At least one	Null	0.68 (0.48 to 0.98)		49.75 (1.60 to 97.91)	

*HR and 95% CI are based on fitting the Cox proportional hazard model (for details see the Methods section).

†Estimate and 95% CI are based on the hierarchical mixed-effects model (for details see the Methods section).

‡Interaction p value was based on a likelihood ratio test by comparing models with and without interaction terms. FEV₁, forced expiratory volume in 1 s.

glutathione, may be important in the interactions with *NOS2A* variants. As no functional data are currently available for the SNPs and the haplotypes studied, detailed sequence analyses followed by mechanistic studies examining different combination of SNPs in the region are needed to identify the specific variants that account for the haplotype associations.

One of the strengths of this study is the prospective follow-up of large numbers of school-age children with annual assessment of asthma diagnosis and lung function measurements in a consistent manner. The associations were robust and highly significant in both PC- and haplotype-based analyses, and the individual SNPs were also significant after adjusting for multiple testing using the p_{ACT} method.³² Furthermore, population admixture is an unlikely explanation of our findings as the incidence rate of new-onset asthma and lung function growth rate during adolescence did not vary by ethnicity and the main effects of the SNPs were similar in the ethnic-specific analysis (online tables E3 and E4). Furthermore, all analyses were adjusted for ancestry factors (based on 233 ancestry informative marker, SNPs) in addition to the traditional self-identified race identifier, thus controlling for any confounding effect of population stratification.

One potential limitation of our study, accuracy of self-reported new-onset asthma assignment, was addressed by excluding any child with a history of wheezing at study entry from the analyses to minimise any major misclassification. A recent study noted that children as young as 7 years of age can provide information regarding their asthma with an acceptable level of validity and reliability.³⁴ Furthermore unless the diagnostic accuracy varied by genotype, error in determining asthma status would probably attenuate the risk estimates and, therefore, would not explain our observed associations. The associations were similar in sensitivity analyses restricted to cases that recently used inhaled medication and were statistically significant for the association between h1000010 and new-onset asthma (online table E8: model 3). Moreover, the observed associations of the promoter SNPs and haplotypes were consistent for lung function growth, which is an objective measurement of respiratory health and not susceptible to diagnostic or reporting bias. Therefore, our results are unlikely to be explained by misclassification of outcome.

We considered the potential effects of selection bias, as genetic data were available for about two-thirds of the initial cohort. As we have described previously,⁴ demographic and socio-economic factors, exposure to maternal smoking during pregnancy and secondhand smoke after birth, and household factors showed modest differences between participants and non-participants. However, adjustment for these factors did not explain our results

(online table E8: model 1), indicating that selection bias based on these or related factors is unlikely to explain our findings.

We conclude that genetic variants of the *NOS2A* promoter region play a role in the respiratory health of children during their adolescence. The role of regulation and function of *NOS2A* expression in asthma pathogenesis and lung function growth and the joint effects with *GSTM1* merit further investigation. Future experimental studies are necessary to define and test variants found by detailed resequencing of the promoter region of *NOS2A* to better understand its role in NO synthesis and respiratory health.

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