Abstract
Background: Asthma is a heterogeneous disease with variable symptoms especially in children. Exhaled nitric oxide (FeNO) has proved to be a marker of inflammation in the airways and has become a substantial part of clinical management of asthmatic children due to its potential to predict possible exacerbation and adjust the dose of inhalant corticosteroids.

Objectives: We analyzed potential factors that contribute to the variability of nitric oxide in various clinical and laboratory conditions.

Material and methods: Study population consisted of 222 asthmatic children and 27 healthy control subjects. All children underwent a panel of tests: fractioned exhaled nitric oxide, exhaled carbon monoxide, asthma control test scoring, blood sampling, skin prick tests, and basic spirometry.

Results: FeNO and other investigated parameters widely changed according to clinical or laboratory characteristics of the tested children. Asthmatics showed increased levels of FeNO, exhaled carbon monoxide, total serum IgE, and higher eosinophilia. Boys had higher FeNO levels than girls. We found a significant positive correlation between FeNO levels and the percentage of blood eosinophils, %predicted of forced vital capacity, total serum IgE levels, and increasing age.

Conclusions: Various phenotypes of children’s asthma are characterized by specific pattern of the results of clinical and laboratory tests. FeNO correlates with total serum IgE, blood eosinophilia, age, and some spirometric parameters with different strength. Therefore, the coexistence of atopy, concomitant allergic rhinitis/rhinoconjunctivitis, and some other parameters should be considered in critical evaluation of FeNO in the management of asthmatic children.

Key words: bronchial asthma, exhaled carbon monoxide, exhaled nitric oxide, children

INTRODUCTION
Asthma is a heterogenous disease with variable signs and symptoms expressed inter- but also intraindividually, especially in children. There are not really specific diagnostic tools or surrogate markers for detecting asthma in infancy. Therefore, diagnosis is usually possible only through long-term follow-up [1]. Another important task is the optimal monitoring of children with confirmed asthma diagnosis and predicting the exacerbations. Beside detailed and repeated analysis of history and careful physical examination, several tests can be applied for this purpose: lung function testing, exercise testing and inflammometry (exhaled nitric oxide, exhaled carbon monoxide, 8-isoprostanes, alkanes, exhaled breath condensate analysis) [1, 2]. Exhaled nitric oxide (FeNO) has proved to be a marker of inflammation in the airways and after initial use for investigation purposes, recently, the measurement of FeNO has become a substantial part of clinical management of asthmatic children due to its potential to predict possible exacerbation and adjust the dose of inhalant corticosteroids. It has been demonstrated that FeNO is elevated in various phenotypes of bronchial asthma (especially in atopic asthma) and that the metabolites of FeNO regulate airway inflammation and airway diameter. It is known that FeNO is not highly specific for pulmonary inflammation, since its can also be observed in non-asthmatic patients suffering from allergic rhinitis [3] or atopic eczema [4, 5]. Many studies searched and analyzed potential factors that regulate production of NO in various pathological and physiological conditions and which are associated with elevated levels in exhaled breath. The evaluation of the degree of asthma control is sometimes difficult, especially in children. For this purpose, FeNO has been introduced into daily practice as an objective parameter assessing, to some extent, the asthma control. FeNO should be evaluated critically, as many factors besides the inflammation presented in the airways modify its variability and levels [6].

In the present study, we aimed to compare differences among various subgroups of asthmatic children in the results of standardized tests and examinations commonly used in management of bronchial asthma. Another important task was to analyze all potential factors that are attributable to changes in fractioned exhaled nitric oxide.
MATERIAL AND METHODS

The study was approved by the Ethical Committee of Jessenius School of Medicine in Martin, Slovakia. Written informed consent was obtained from the parents of all tested children. We examined a population of 249 schoolchildren (aged 12 ± 4 years, 55% boys). The study was performed at the Department of Pediatrics in University Hospital in Martin, Slovakia. Bronchial asthma was defined as recurrent airways obstruction manifested by wheeze and dyspnea relieved spontaneously or by bronchodilator therapy, as defined in GINA. Altogether, we enrolled 222 asthmatic children (89%) and 27 healthy control subjects of comparable age without any allergic disease or subclinical atopy (11%). None of the studied subjects was smoking. The children or their parents filled standardized Asthma control test® and questionnaire aimed at personal and family history of allergic and respiratory diseases, frequency of upper or lower respiratory infections, presence of pets at home, actual clinical complaints and signs of respiratory disease, and provided therapy. The children underwent blood sampling for blood cell count and basic biochemical analysis (C-reactive protein, total IgE in serum) and also the measurement of FeNO and exhaled carbon monoxide (eCO). FeNO was measured online according to the recommendations of the ERS/ATS standards [7], using a portable nitric oxide analyzer (NIOX-MINO®, Aerocrine, Sweden). eCO was measured with a Micro 4 Smokerlyzer® (Bedfont, England). We also performed basic spirometry (KoKo DigiDoser-Spirometer; nSpire Health, Louisville, CO) according to the ATS/ERS recommendations [8]. The FeNO and eCO values were estimated prior to the spirometric examination.

Due to the possibility of discontinuance of antihistamine treatment, we performed a skin prick tests (SPT), with a panel of common inhalant allergens (Dermatophagoides pteronyssinus; cat dander; dog dander; feather mix; mixed 4 cereals; mixed 5 grasses: Avena sativa, Hordeum vulgare, Secale cereale, Triticum sativum; mixed moulds: Cladosporium herbarum, Aspergillus mix, Alternaria alternata; Artemisia vulgaris, Artemisia folia, Betula alba) (ALK-ABELLO, Horsholm, Denmark), and we measured the levels of specific serum IgE (sIgE) against the same allergens (ELISA). The SPT with allergens was defined as positive if the wheal was ≥3 mm in its longest dimension. All the tests were performed by the same well-trained operator. The children with at least one positive SPT or sIgE were considered as atopic. Data are expressed as means ±SE unless otherwise indicated and were analyzed with the SPSS version 9.0 (SPSS Inc. Chicago, IL). Student’s two-tailed t-test or Mann-Whitney U non-parametric test was used for statistical comparison. Spearman’s correlation (rS) was used for correlation studies. P ≤0.05 was considered to indicate statistical significance.

RESULTS

Regarding the clinical control of asthma, 28 children (12%) underwent testing during acute exacerbation (shortness of breath, wheezing, chest tightness, and/or coughing), while the rest of the children (87%, 194/222) had clinically controlled asthma. Thirty percent of the asthmatics (67/222) did not suffer from any other allergic disease, but 70% of the children (155/222) also had symptoms of concomitant allergic rhinitis/rhinoconjunctivitis (ARC). According to the results of SPT and specific IgE, 69% of the children suffered from atopic asthma (126/183) and the rest was non-atopic (57/183, 31%). Non-compliance with recommended therapy was noted in 23% of the children.

FeNO and other investigated parameters substantially differed according to clinical or laboratory characteristics of the tested children (Table 1). Comparing asthmatic children with healthy subjects, we found increased FeNO (25.2 ± 1.8 vs. 12.4 ± 1.2 ppb, P=0.043) and eCO (1.0 ± 0.1 vs. 0.4 ± 0.2 ppm, P=0.013). Asthmatics had a higher level of eosinophilia in peripheral blood (absolute: 0.41 ± 0.12 x 10³ vs. 0.14 ± 0.02 x 10³/ml, P=0.001; percentage: 4.0 ± 0.3 vs. 2.2 ± 0.3%, P=0.005) and of total serum IgE (234.4 ± 26.3 vs. 31.8 ± 4.8 U/ml, P=0.005) compared with controls. All spirometric parameters were lower in asthmatics.

Table 1. Comparison of selected parameters in subgroups of asthmatic children and healthy controls.

|                  | FeNO (ppb) | eCO (ppm) | ACT (score) | Serum IgE (U/ml) | Eosinophils (10⁹/l) | Eosinophils (%) |
|------------------|------------|-----------|-------------|------------------|----------------------|----------------|
| BA               | 25.2±1.8   | 1.0±0.1   | 22.7±0.2   | 234.4±46.3       | 0.41±0.12            | 4.0±0.3        |
| BA acute         | 63.0±8.6***†† †† | 1.2±0.2†   | 20.6±0.8**†  | 503.2±136.7***††† | 0.46±0.15†††        | 5.2±1.4†        |
| BA stable        | 20.1±1.3   | 1.0±0.1†† | 22.9±0.2   | 200.5±23.2†      | 0.40±0.12†‡          | 3.8±0.2‡        |
| BA atopic        | 31.2±2.8***††† | 1.0±0.1†† | 22.6±0.3   | 285.9±41.1***††† | 0.52±0.18***†††     | 4.8±0.4***†††   |
| BA non-atopic    | 14.9±1.6   | 1.1±0.1†   | 23.4±0.3   | 115.8±26.4       | 0.28±0.10†          | 2.9±0.3         |
| BA isolated      | 20.5±2.7   | 0.9±0.1‡   | 22.8±0.4   | 204.2±20.4†      | 0.28±0.05†          | 3.8±0.5‡‡       |
| BA + ARC         | 27.3±2.3***††† | 1.1±0.1†| 22.6±0.3   | 247.3±33.5†       | 0.47±0.16†††        | 4.1±0.3†        |
| Controls         | 12.4±1.2   | 0.4±0.2    | ---        | 31.8±4.8         | 0.14±0.02            | 2.2±0.3         |

FeNO – fractioned exhaled nitric oxide, eCO – exhaled carbon monoxide, ACT – asthma control test, BA – bronchial asthma, ARC – allergic rhinoconjunctivitis; Comparison between two asthmatic subgroups: *P<0.05, **P<0.01, ***P<0.001; Comparison between asthmatic subgroup and healthy controls: †P<0.05, ††P<0.01, †††P<0.001.
but the differences did not reach statistical significance. Analyzing the results among the groups of children divided according to the clinical control, the highest levels of FeNO were in asthmatic children during acute exacerbation compared with both stable asthmatics (63.0 ± 8.6 vs. 20.1 ± 1.3 ppb, *P* <0.001) and healthy controls (63.0 ± 8.6 vs. 12.9 ± 1.4 ppb, *P* <0.001). Stable asthmatics did not differ with controls regarding the FeNO, eCO, total serum IgE, and eosinophils in peripheral blood were in general higher in asthmatics compared with healthy children, but we did not find the difference in eCO between asthmatic in acute and stable phases of the disease. Acute asthmatics had a lower total score in asthma control test than stable asthmatic children (20.6 ± 0.8 vs. 22.9 ± 0.2, *P* = 0.002). Neutrophils in blood were increased in acute (absolute: 5.69 ± 0.63 x10⁹ vs. 2.06 ± 0.32 x 10⁹/l, *P* = 0.035), but not in stable asthmatics compared with controls. We found similar results in the analysis of atopic and non-atopic asthma.

Atopic and non-atopic asthmatics did not differ in eCO and total scores in asthma control test. We observed higher FeNO, total serum IgE, and blood eosinophils in atopic asthmatics in comparison with non-atopic and healthy asthma and control children. Interestingly, while comparing acute and stable asthmatics, atopic and non-atopic asthmatics between each other and with healthy subjects, we found only slightly decreased spirometric parameters in asthmatics, but comparing the children with isolated form of asthma with those who suffered from concomitant allergic rhinitis/rhinoconjunctivitis, several significant differences appeared. Isolated asthma was characterized by a significant reduction in FEV1 and FVC (compared to children with asthma+ARC) and with the decrease of FEF25-75% and PEFR25-75% (compared with both children with asthma+ARC and healthy controls). Regarding the atopy status, 63% cases of isolated asthma and 71% of asthma associated with ARC were atopic (n.s.). Atopic asthmatics with concomitant ARC did not differ in any parameter compared with healthy children. Asthma without ARC showed the same values of FeNO and eCO as did healthy controls, but differed in total serum IgE and blood eosinophils. Both groups of asthmatic children showed no differences in eCO, asthma control test, total serum IgE levels, or blood eosinophils (absolute and percentage). Asthmatics with concomitant ARC had elevated of FeNO, eCO, total serum IgE, and blood eosinophils compared with healthy controls and higher FeNO than isolated asthma (Table 1). Children with non-compliance to treatment had higher FeNO (46.4 ± 4.9 vs. 18.1 ± 1.3 ppb, *P* = 0.001) and total serum IgE (371.0 ± 73.0 vs. 172.7 ± 23.0 U/ml, *P* = 0.010) compared with those who accepted our recommended therapy. Subjects with positive family history for allergic diseases (bronchial asthma, allergic rhinitis/rhinoconjunctivitis, atopic eczema) showed higher FeNO (27.8 ± 2.8 vs. 20.1 ± 1.8 ppb, *P* = 0.067) and blood eosinophils (absolute: 0.56 ± 0.20 x10⁹ vs. 0.21 ± 0.02 x 10⁹/l, *P* <0.001; percentage: 4.6 ± 0.4 vs. 3.0 ± 0.3%, *P* = 0.035). Regarding the gender, boys had higher FeNO than girls (20.9 ± 2.5 vs. 20.6 ± 2.0 ppb, *P* = 0.048), eCO (1.2 ± 0.1 vs. 0.7 ± 0.1 ppm, *P* = 0.002), but did not differ in total serum IgE or eosinophil levels.

We also analyzed the possible correlation between the FeNO values and the results of other laboratory and clinical examinations. FeNO levels correlated significantly with the percentage of blood eosinophils
(Fig. 1A), total serum IgE levels (Fig. 1B), %predicted forced vital capacity (Fig. 1C), and increasing age of the tested subjects (Fig. 1D). We found a marginally significant negative association between FeNO and the total score of asthma control test (not shown). FeNO was not related to eCO levels.

**DISCUSSION**

We analyzed differences in various indices measured in several subgroups of asthmatic children and healthy controls. We divided the children according to the differing characteristics: asthmatics vs. healthy control children, exacerbated vs. stable asthmatics, atopic vs. non-atopic children and those with isolated allergic rhinitis/rhinoconjunctivitis. The indices widely differed according to the clinical or laboratory characteristics of tested asthmatics. The levels of FeNO were increased in asthmatics in general, during acute exacerbation of the disease, in asthmatics with atopy and in children with asthma and other respiratory allergic co-morbidity. Interestingly, we were unable to detect significant differences in spirometric indices between asthmatics and healthy controls. It has been shown that the measurement of FeNO is more reliable in the diagnostic algorithm of BA than the results of lung function testing [9, 10]. The application of FeNO as a diagnostic test for asthma could be limited by lower sensitivity. However, good specificity of FeNO could, in the presence of its low level, exclude the diagnosis of BA [11]. It has been suggested that elevated FeNO may help alert the clinician to the possibility of undiagnosed asthma [12]. Although none of the developed methods is optimal to predict an exacerbation of asthma, it has been confirmed that FeNO is helpful to this end, and it also could be used to guide therapy, adjust inhaled corticosteroids dose, or predict [13]. Smith et al [9] demonstrated that when comparing standard guidelines-directed care to the addition of FeNO, the latter group showed a significant reduction in the maintenance therapy required, while simultaneously achieving a trend toward a decrease in asthma exacerbations. The results of their study supported the hypothesis that control of airway inflammation can improve asthma control [9]. A combination of FeNO and FEV1 results could increase the sensitivity of FeNO in predicting the relapse of asthma [14]. It is confirmed by many authors that acute respiratory infections, which are usually the cause of asthma exacerbation, are connected with concomitant increases in FeNO. Interestingly, however, Kovesi et al [12] found that healthy children of general, unselected population without asthma, but with reported bronchitis or pneumonia in the past year had abnormally high FeNO concentrations without the presence of asthma. In the present study, asthmatics who did not comply with therapy had increased FeNO, which suggests that this parameter could also be used as an indicator of non-following the therapeutic recommendations [15]. Similarly to our findings, children with asthma alone had lower FeNO than children with asthma and other allergies (eczema, rhinitis, and rhinoconjunctivitis) [12]. Welsh et al [16] found in a multivariate analysis that asthma, hay fever, and atopic dermatitis all significantly influenced FeNO in children. Thus, elevated FeNO could also result from atopic disease other than asthma [12]. Healthy children with positive allergy skin tests with or without serum eosinophilia may have elevated FeNO regardless of whether symptoms of asthma or allergic rhinitis are present [17]. Atopic asthma is associated with an elevation of FeNO in comparison with subjects suffering from non-atopic variant of this disease [18]. Recent study has revealed that both allergic and non-allergic asthma are related to increased FeNO levels, but only in never-smoking subjects. A limited value of FeNO to detect subjects with asthma among ex- and current smokers suggests the predominance of a non-eosinophilic inflammatory phenotype of asthma among never-smokers [19].

Isolated asthma did not differ in eCO from controls, but asthmatic children with allergic rhinitis showed elevation of eCO. Increased eCO in asthmatics may reflect the expression and activation of heme oxygenase-1 in airways, due to inflammation and oxidative stress [20]. eCO concentration is increased during early and late phase of asthmatic reaction to allergen exposure, whereas FeNO is increased only in the late phase [21]. Although the differences in eCO are not as expressive as those in FeNO, elevation of eCO correlates not only with airway inflammation, as does FeNO, but also with increased oxidative stress in asthmatic airways [2]. eCO could add information in the management of bronchial asthma, and also can help detect smokers among adolescent asthmatics, which, in turn, can help in critical evaluation of FeNO levels obtained in these patients.

Another aim of the present study was to analyze other potential factors that may influence FeNO. Elevated FeNO was associated with higher total serum IgE levels and eosinophilia, FVC, and increasing age. Boys had higher FeNO than girls. Other authors also found a significant contribution of gender [5, 6, 12], age [5, 6, 22], serum total IgE [17, 22, 23, 24], or eosinophils [17, 22]. In atopic asthmatics, serum IgE levels appear to reflect systemic (blood eosinophils) and organ-specific (FeNO) markers of allergic inflammation [22]. While we noted a weak but significant correlation between FeNO and some spirometric indices, other studies did not confirm that relationship [12]. Borderline association between the asthma control test score and FeNO suggests that these two parameters describe two different aspects of asthma – clinical symptoms and inflammation in airways [25]. Since the correlation coefficients were in some cases low, although significant, it is probable that also other intrinsic and extrinsic factors significantly contribute to the FeNO variability, such as smoking, atopic status, concomitant upper or lower respiratory infection, testing during pollen season, allergen content in the air, presence of atopic eczema, etc. [11, 12, 17].

We conclude that FeNO could not be expected to be the perfect inflammometer. However, FeNO presents an easy test which is helpful to target steroid treatment and to provide significant benefits in terms of less hyperresponsiveness and less inflammation. FeNO offers more for the day-to-day asthma management than any of the conventional lung function tests.
The variability of FeNO results reported in the present study suggests the possible interplay of various bioactive factors in the formation and release of NO, which should be taken into account in clinical decision-making based on the FeNO measurement.

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Corresponding author: Milos Jesenak, M.D., Ph.D., MBA Department of Pediatrics Jesseniess School of Medicine Kollarova 2 str. 036 59 Martin, Slovakia Phone: +421 903 275331 Fax: +421 043 4222678 jesenak@gmail.com

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