Exhaled Breath Temperature in Asthmatics and Controls after Eucapnic Voluntary Hyperventilation and a Methacholine Challenge Test

Henning Svensson  Leif Bjermer  Ellen Tufvesson
Respiratory Medicine and Allergology, Department of Clinical Sciences, Lund University, Lund, Sweden

Key Words
Asthma · Eucapnic voluntary hyperventilation · Methacholine · Exhaled breath temperature

Abstract
Background: It has been suggested that exhaled breath temperature (EBT) is increased in asthmatic subjects. Objectives: Our aim was to investigate EBT in asthmatics compared to healthy controls before and after eucapnic voluntary hyperventilation (EVH) and a methacholine challenge test (MCT). Methods: A total of 26 asthmatics and 29 healthy controls were included. Forced expiratory volume in 1 s (FEV₁), EBT and oral, axillary and auricular temperatures were measured before and after EVH and MCT. Results: FEV₁ % predicted (%p) was significantly lower in asthmatic subjects compared to healthy controls at all time points. EBT was significantly increased in all subjects 15–30 min after EVH and 5–45 min after MCT. Oral temperature displayed a similar pattern of increase, in contrast to axillary and auricular temperature, and correlated with EBT before and after both of the challenge tests. EBT after 5 min correlated with the largest drop in FEV₁%p after EVH in asthmatic subjects. No significant differences or changes in EBT were found when comparing asthmatics to healthy controls before or after any of the tests. Conclusions: EBT is increased after both EVH and MCT, possibly reflecting a vascular response. This is related to both the fall in FEV₁ and to oral temperature, suggesting an effect on the whole respiratory tract including the oral cavity. No differences in EBT are seen between asthmatics and healthy controls, indicating that the increase in EBT is mainly physiological rather than pathophysiological.

Introduction
Asthma is characterized by inflammation of the airways, with inflammatory cells present as well as airway remodeling, increased vascularization and bronchial hyperresponsiveness (BHR). Provocation testing for detecting BHR in asthmatics has gained importance for diagnosis and for monitoring disease activity, guiding the dosing of inhaled steroids [1].

Exhaled breath temperature (EBT) is a potential biomarker for airway inflammation. It has been shown to relate to disease exacerbations [2–4] as well as to vascular [5] and fibrous remodeling [6] of the airways, and to correlate with levels of exhaled nitric oxide (NO) in childhood and adult asthma [7, 8]. Sub-epithelial blood vessel density is higher in asthmatic subjects than in healthy controls [9], and the hypothesis is that increased blood flow during airway inflammation would result in heating of the ventilated air to a higher temperature [5]. Some studies have shown that EBT is elevated in asthmatics compared to healthy controls [2, 8, 10], but research on the relationship between EBT and BHR is limited. In a previous study, we found that an increase in EBT corre-
lated with a decrease in forced expiratory volume in 1 s (FEV₁) in both asthmatics and controls after an exercise challenge test [11]. A similar correlation has been found in asthmatic children [12]. In addition, we found that there was a prolonged increase in EBT after exercise in asthmatics for whom the FEV₁ fell during the test compared to asthmatic subjects for whom it did not fall. This indicates a link between EBT and airway inflammation, but the potential of a clinical implication of EBT is yet to be determined. Increased baseline levels of 8-isoprostane, a marker of oxidative stress, have been found in exhaled breath condensate of asthmatic children with BHR after an exercise challenge test, supporting the notion that persistent airway inflammation is important for exercise-induced bronchoconstriction to occur [13].

Eucapnic voluntary hyperventilation (EVH) is an indirect challenge test, similar to the exercise challenge test, where the inflammatory cells of the airways are triggered to release mediators [14, 15], causing smooth muscle cell constriction and vasodilatation. The triggering factor for this release is believed to be an increased osmolarity of the periciliary fluid covering the respiratory mucosal membrane [16], due to increased loss of water to the air when there is a high rate of ventilation. The specificity of an indirect challenge test is considered high, as, in theory, inflammation is required for a positive result to occur.

Methacholine is believed to act directly on the mucociliary receptors [17] of smooth muscle cells, endothelial cells and mucus-producing cells, inducing bronchial obstruction, vasodilatation and mucus production. The sensitivity of a methacholine challenge test (MCT) is high, while a positive test has a low diagnostic specificity. The presence of inflammation and chronic remodeling of the airways are both believed to increase reactivity to methacholine [18].

The aim of this study was to investigate changes in EBT after EVH and MCT, and to compare it to changes in body temperature. EVH is believed to trigger bronchoconstriction and vasodilatation through the release of inflammatory mediators, similar to an exercise challenge test. Methacholine was used as an additional challenge, reflecting another pathophysiological mechanism. We hypothesized that those subjects who responded to EVH with bronchoconstriction would increase their EBT to a greater extent than those who did not, due to the probable vasodilatation following the release of mediators from airway inflammatory cells. An increase in EBT would also theoretically be seen after MCT, where the pharmacological effect of methacholine acts not only on smooth muscle cells but also induces vasodilatation. A secondary aim was to investigate whether an increase in airway resistance would affect the change in EBT, due to narrowing of the airways.

### Materials and Methods

#### Subjects

Twenty-six subjects with a doctor’s diagnosis of asthma according to the guidelines of the Global Initiative for Asthma [19] were investigated (table 1). Twenty-nine healthy subjects with no diagnosis of asthma or any respiratory symptoms were used as controls. All subjects were interviewed concerning their previous and present health, and asthmatic subjects filled out an Asthma Control Test questionnaire. Subjects with respiratory tract infections (within the last 3 weeks) or any other medical conditions (apart from asthma) affecting their health were excluded, as were subjects with a history of smoking. The subjects were not allowed to drink coffee for at least 6 h prior to either challenge test or to conduct any form of exercise on the same day. All asthmatic subjects refrained from using short-acting β2-agonists for at least 8 h and long-acting β2-agonists and inhaled corticosteroids (ICS) for at least 24 h prior to any part of the study. All subjects gave written informed consent and the study was approved by the regional Ethics Review Board, Lund.

#### Study Design

We conducted tests from the 14 July 2011 to the 1 August 2012. Two different respiratory tract provocation tests, EVH and MCT, were performed in random order by all subjects. At least 48 h passed between tests. Prior to each test, and after at least 5 min of rest, exhaled NO, spirometry, EBT and axillary, auricular and oral temperatures were measured (in the given order). Thereafter, the subjects performed a provocation test. Afterwards, measurements of spirometry, EBT, axillary, auricular and oral temperatures were

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**Table 1. Subject characteristics**

<table>
<thead>
<tr>
<th></th>
<th>Controls (n = 29)</th>
<th>Asthmatics (n = 26)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Males/females, n</td>
<td>16/13</td>
<td>16/10</td>
</tr>
<tr>
<td>Age, years</td>
<td>25 (23–26)</td>
<td>24 (22–27)</td>
</tr>
<tr>
<td>Baseline FEV₁, %p</td>
<td>104 (97–107)</td>
<td>95 (90–98)</td>
</tr>
<tr>
<td>FEV₁ drop ≥10% after EVH, n percentage units</td>
<td>6 (4–9)</td>
<td>10 (6–15)</td>
</tr>
<tr>
<td>FEV₁ drop ≥20% after MCT, n percentage units</td>
<td>7</td>
<td>16</td>
</tr>
<tr>
<td>Fall in FEV₁, %p after EVH, percentage units</td>
<td>10 (7–20)</td>
<td>21 (13–28)</td>
</tr>
<tr>
<td>Atopy, n</td>
<td>7</td>
<td>20</td>
</tr>
<tr>
<td>ICS (400–800 μg budesonide/day), n</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>Exhaled NO, ppb</td>
<td>14.7 (9.9–23.5)</td>
<td>17.1 (14.9–26.4)</td>
</tr>
</tbody>
</table>

Data are expressed as a median (interquartile range), where appropriate. For baseline FEV₁, % and exhaled NO values, a mean of the two registrations before each test is used for each subject.
performed repeatedly during a period of rest, as described below. After completing the measurements, 60 min after the provocation test, 400 µg of salbutamol was inhaled by the subject. After another 15 min of rest, a final set of measurements (spirometry, EBT and body temperatures) was conducted.

**Eucapnic Voluntary Hyperventilation**

The subjects hyperventilated for 6 min through a mouthpiece connected to a container administering dry hypercapnic air (5% CO₂) via a dry air ventilation device (Aiolos Medical AB, Karlstad, Sweden). The air was at an ambient temperature. A nose clip was used, and the flow was set to 26 × FEV₁ (baseline value) l/min. The subjects were supervised at all times and guided by a reservoir balloon attached to the ventilation device, ensuring that the rate and depth of breathing were adequate. The test was considered positive if FEV₁ dropped by ≥10% (compared to the baseline value) at any measurement within 30 min of the end of the hyperventilation.

**Methacholine Challenge Test**

For the MCT, a tidal-volume-triggered device (Aerosol Provocation System, APS; Erich Jaeger GmbH) was used. One inhalation of 9 mg/ml NaCl was performed first as a negative control. Methacholine was then inhaled repeatedly, with a spirometry performed 2 min after each inhalation and the next inhalation following directly thereafter. Five inhalations were administered with increasing doses (50, 150, 300, 600 and 900 µg), resulting in a maximal cumulative dose of 2,000 µg of methacholine. A positive test was defined as a drop of FEV₁ by ≥20% to baseline value, in which case the challenge test was considered completed and measurements were made as described below.

**Exhaled NO Measurement**

Exhaled NO was measured prior to each of the tests using a NIOX Flex (Aerocrine AB, Stockholm, Sweden) according to recommendations of the American Thoracic Society and European Respiratory Society [20] with an exhalation flow rate of 50 ml/s.

**Lung Function Test**

Flow volume spirometry (Jaeger MasterScope, Würzburg, Germany) measuring FEV₁ was performed before and at 5, 10, 15, 20, 30, 45 and 60 min after the end of each of the provocation tests, and 15 min after inhaling salbutamol. FEV₁ % predicted (%p) was calculated according to the reference spirometric values of Crapo et al. [21].

**EBT Measurement**

EBT was measured using an X-Halo (Delmedica Investments, Singapore) once before and at 5, 15, 30, 45 and 60 min after each of the tests, as well as 15 min after inhaling salbutamol. Subjects were instructed to breathe tidally, inhaling through the nose and exhaling through the mouth into the device. Measurements took 2–6 min to complete.

**Body Temperature Measurements**

Body temperatures were measured before and at 5, 15, 30, 45 and 60 min after each of the provocation tests as well as 15 min after inhaling salbutamol. Oral and axillary temperatures were measured with digital clinical thermometers C402 and C202, respectively (Terumo, Leuven, Belgium). Auricular temperature was measured using a ThermoScan type 6013 (Braun, Kronberg, Germany).

**Impulse Oscillometry System**

A Jaeger MasterScreen impulse oscillometry system (IOS; Erich Jaeger GmbH) was used to test a subgroup consisting of 22 of the subjects (9 asthmatics and 13 healthy controls). IOS was performed before MCT and after each inhalation during the challenge prior to spirometry, as previously described [22]. Subjects used a nose clip and were instructed to press the palms of their hands against their cheeks in order to decrease the influence of movements of the upper airways on IOS parameters. Oscillometric pressure impulses with a pulse sequence of 5 per second and a frequency spectrum between 5 and 35 Hz were superimposed on tidal breathing for at least 30 s. Airway resistance at 5 Hz (R5) and 20 Hz (R20), reactance at 5 Hz (X5), resonant frequency (Fres) and area of reactance integrated from 5 Hz to Fres (AX) were determined, and R5–R20 was calculated.

**Statistical Analyses**

GraphPad Prism version 4.0 was used for statistical analysis. The results are expressed as median (interquartile range) where appropriate. Correlations were calculated using Spearman’s correlation test. Comparisons between groups were made by using the Mann-Whitney test. Analysis of paired measurements was made by using Wilcoxon’s test. A p value of <0.05 (two-tailed) was considered significant. Power calculation was performed based on 2 independent groups of 26 evaluable subjects. With an expected standard deviation of 0.5, it was possible to detect a significant difference with a power of 94%.

**Results**

**Subjects**

Twenty subjects (14 asthmatics and 6 controls) had a positive EVH test and 23 (16 asthmatics and 7 controls) had a positive MCT (table 1). Seventeen of the subjects (12 asthmatics and 5 controls) were positive for both challenge tests. There were no significant differences in EBT or ΔEBT (defined as the difference between baseline EBT and the highest EBT after the challenge test) when comparing those who were positive for both challenge tests to those who were positive for one test only or negative for both tests. The Asthma Control Test score among asthmatic subjects was 21.5 (20–23), and it did not differ between subjects who had a positive EVH or MCT and those who did not.

**Lung Function**

As expected, levels of FEV₁ %p were significantly lower in asthmatic subjects than controls at baseline (p = 0.002–0.006) and 5–60 (p = 0.001–0.006) min after both tests, and also after inhaling salbutamol (p = 0.003–0.007).

All subjects (both asthmatics and controls) experienced a drop in the level of FEV₁ after both EVH and MCT (p < 0.001), and FEV₁ increased postsalbutamol...
compared to at both baseline (p < 0.001 after EVH; p = 0.040 after MCT) and 60 min after the challenge tests (p < 0.001). A majority of the subjects had their greatest drop in FEV₁ at 5–10 min after both EVH (n = 34) and MCT (n = 42). FEV₁% in all subjects was lowest 5 min after EVH [90.5% (83.4–100.2)] and 5 min after MCT [83.4% (74.7–100.4)].

During EVH, FEV₁% was significantly lower at all time points (except for 45 min) in asthmatic subjects with ICS treatment than in those not treated with ICS. Following MCT, FEV₁% was lower in asthmatic subjects treated with ICS after 15 min and postsalbutamol, but not at any other time point. There was no difference in FEV₁% between male and female subjects.

**Exhaled-Breath Temperature**

EBT was significantly increased compared to baseline following EVH in all subjects after 15 min, and it peaked at 30 min (p < 0.001; fig. 1a). After MCT, EBT was significantly increased in all subjects compared to baseline at 5–45 min (p < 0.001–0.006; fig. 2a). EBT peaked 15 min after MCT.

There were no significant differences between asthmatics and healthy controls in EBT or ΔEBT at any time point during the challenge tests (fig. 3) or between subjects with and without a positive challenge test. There were no differences in baseline EBT or ΔEBT between asthmatic subjects with and without ICS treatment, between atopic and nonatopic subjects or between male and female subjects.

**EBT Correlation with Lung Function**

ΔFEV₁% (defined as the maximum decrease in FEV₁%: baseline FEV₁% – lowest FEV₁% after the challenge test) after EVH correlated with EBT measured after 5 min in asthmatic subjects (p = 0.048, r = 0.392), but not in controls (p = 0.641, r = –0.091), all subjects (p = 0.421, r = 0.111; fig. 4a) or when examining into subjects with a positive or negative EVH challenge test (data not shown).

No similar correlation between ΔFEV₁% and EBT 5 min was seen after MCT in the asthmatics, the controls or all subjects (fig. 4b).

There was no correlation between ΔFEV₁% and ΔEBT after any of the challenge tests.
Body Temperatures and Correlations with EBT

Oral temperature displayed an increase after both EVH and MCT, similar to what was seen in EBT. Oral temperature was significantly elevated in all subjects 30 min after EVH (p = 0.003; fig. 1b). After MCT, oral temperature was elevated compared to baseline after 5–30 min (p < 0.001–0.046; fig. 2b).

Neither axillary nor auricular temperatures showed a similar pattern of increase after EVH (fig. 1). On the contrary, axillary temperature was decreased and auricular
temperature was not affected. After MCT, auricular temperature was increased at 5 min (p < 0.001) and 15 min (p = 0.013) compared to baseline value (fig. 2d), while axillary temperature instead dropped after 45 min (p = 0.020) and 60 min (p = 0.009; fig. 2c).

EBT correlated with oral temperature at baseline (p = 0.044, r = 0.392), before EVH and 5 min (p = 0.032, r = 0.288), 45 min (p = 0.005, r = 0.372) and 60 min (p = 0.029, r = 0.292) afterwards as well as postsalbutamol (p < 0.001, r = 0.501). EBT and oral temperature also correlated before and after MCT at baseline (p = 0.003, r = 0.390) and after 15–60 min (p = 0.001–0.015, r = 0.325–0.424). Peak oral temperature coincided with peak EBT after both EVH (fig. 1) and MCT (fig. 2).

ΔEBT and Δoral temperature (defined as the difference between baseline oral temperature and maximum oral temperature after the challenge test) correlated significantly after EVH (p = 0.001, r = 0.428) and MCT (p = 0.014, r = 0.330; fig. 5). No corresponding correlations were seen when comparing ΔEBT to axillary or auricular temperature (data not shown).

Exhaled Nitric Oxide

Baseline levels of exhaled NO did not differ between asthmatic subjects and controls, between atopic subjects and nonatopic subjects, between asthmatic subjects with or without a positive challenge test or between asthmatic subjects with or without ICS treatment. Levels of exhaled NO did not correlate with EBT before or after any of the challenge tests except postsalbutamol after MCT (p = 0.041, r = 0.277). It did not correlate with ΔEBT.
No significant correlations were found between EBT (including ΔEBT) and IOS parameters at any time point, or between EBT and the change in any IOS parameter compared to at baseline.

As expected, increases in R₅, R₂₀, R₅-R₂₀ and AX, and a decrease in X₅, were seen in all subjects when comparing baseline values to those measured after MCT. FRES was increased in controls only. R₅-R₂₀%p was significantly higher in asthmatic subjects than in healthy controls after MCT (data not shown), but no other IOS parameters differed at baseline or after MCT.

**Discussion**

This study showed that EBT increased significantly 15–30 min after EVH and 5–45 min after MCT, and that no difference in EBT between asthmatic subjects and healthy controls could be detected. Levels of FEV₁%p were significantly lower in asthmatic subjects than in controls at all time points; this confirmed that the selection of subjects was adequate. Furthermore, no difference in EBT was seen when comparing subjects with and without BHR after any of the challenge tests.

The increase in EBT seen after EVH and MCT may be a result of increased blood flow following vasodilatation. However, vascular tone alone does not explain the changes in EBT, seeing as EBT decreased from the time point of 60 min to postsalbutamol after both challenge tests, even though salbutamol is known to increase bronchial blood flow [23]. Other factors such as bronchodilation may also affect EBT, possibly masking the effects of increased blood flow on EBT. Similar studies should be undertaken with the aim of assessing the effects of EVH and MCT on markers of inflammation in exhaled breath condensate, including metabolites [24, 25], leukotriene B₄ [26] and isoprostanes [27, 28] in patients with asthma. Likewise, it would be worth studying the effects of these challenges on e-nose breathprints [29, 30].

A significant increase in EBT was seen already at 5 min after MCT, in contrast to at 15 min after the end of hyperventilation. Inhalation of methacholine affects vascular tone and increases bronchial blood flow [18], possibly leading to an early response in EBT. Increased blood flow after EVH is dependent on dehydration of the periciliary fluid and on the release of inflammatory mediators, which might delay the reaction somewhat and give rise to a relatively mild response in both EBT and decrease of lung function. EBT was elevated for a longer period of time following MCT compared to EVH. The duration of an elevation of EBT after airway provocation tests might be related to the density of the challenge and the airway response. For example, EBT was elevated for an even greater period of time (at least 60 min) following a standardized exercise challenge test [11]. A greater increase in body temperature in general after exercise would not form a satisfactory explanation for this difference, since EBT was elevated independently of the axillary and auricular temperature. The maximum response to exercise (peak EBT) was greater than that after EVH, in spite of the fact that the respective durations of hyperventilation in the two challenge tests are comparable. The increase in cardiac output, with a possible effect on blood flow of the airway mucosa during and after exercise, may provide an explanation for these differences. During EVH, heart rate increases only marginally and blood flow would therefore not be increased to the same extent. Another possibility is that a standardized exercise challenge is a better method of ensuring that ventilation, with subsequent dehydration of the airways, is adequate. While subjects performing EVH may subtly decrease their rate of ventilation to a level perceived as more comfortable, a standardized exercise challenge test performed on a treadmill at approximately 90% of maximum heart rate ensures that the respiratory rate increases considerably.

In this study, we showed that the maximum change in FEV₁%p correlated with EBT in asthmatic subjects 5 min after EVH, but not in the controls. A similar pattern was seen 5 min after the exercise challenge test [11] in all subjects and in the controls, but not when looking at asthmatics separately. The similarities of correlations between EBT and decrease in lung function support the theory that EVH is equivalent to an exercise challenge test, at least in this perspective. However, while EBT reached its peak after 5 min following the exercise challenge test, maximum median EBT after EVH was seen after 30 min.

The maximum increase in oral temperature correlated with a maximum increase in EBT after both EVH and MCT. Oral temperature displayed the same pattern as EBT after both the provocation tests that we used as well as after exercise, proving that the two temperature measurements are closely related. Although the X-Halo has been validated in previous studies [2, 4], one must take into consideration the possibility that since measurements reflect the plateau of the breath temperatures of several expirations registered during a time period of a few minutes, oral temperature in itself may have some effect on the result. This would be in contrast to the single-breath method described in other studies [5–8].
lar temperature increased 5–15 min after MCT, which may possibly be explained by a spread of heat to the ear through the Eustachian tube. However, peak auricular temperature did not coincide with the peaks of EBT and oral temperature after MCT. Axillary temperature was decreased after EVH and to a lesser extent after MCT. The explanation for this is not known, but it is possible that the time of rest following arrival at the laboratory was insufficient, resulting in an elevated axillary temperature at baseline. However, in this case, baseline axillary temperature before MCT would most probably also have been affected.

In summary, our findings suggest that EBT is directly related to oral but not to axillary or auricular temperature, regardless of the type of challenge test or level of physical activity. We believe that this connection is a result of increased blood flow due to vasodilatation taking place only in the airways and/or the oral cavity, both being part of the respiratory tract.

Some of the controls in our study displayed BHR after one or both of the airway challenge tests. The prevalence of positive methacholine challenge tests in the general population has been reported to range from <10% to >40% in various studies, and normal variations in BHR can be expected even among subjects selected for the absence of known potential causes of a positive challenge test, such as asthma [31]. Similar variations are most probable even for indirect tests of BHR, even though they normally display a higher specificity [32].

Exhaled NO has been shown to correlate with EBT in some studies [7, 8], but this was not one of our findings. We also found no differences in levels of exhaled NO between asthmatics and controls, between asthmatics with and without ICS treatment or between atopic and non-atopic subjects. As seen by the results of the Asthma Control Test, this study used subjects with relatively mild and well-controlled asthma, possibly representing a different phenotype from that used in studies showing elevated levels of NO in asthmatics. The fact that there was a significant difference in FEV1%p at all times should nevertheless confirm that our selection of asthmatic subjects was representative. EBT has also recently been suggested to correlate with body height [33], but no such correlations were found in our study.

One hypothesis was that airway narrowing, measured as airway resistance by IOS, would affect changes in EBT. However, no such correlations could be seen.

To conclude our findings, we have shown an increase in EBT after both EVH and MCT, reflecting a normal physiological response which is similar in asthmatic subjects and healthy controls. EBT is probably affected by vasodilating agents and increases when lung function declines, although significant correlations between the two are limited. EBT and oral temperature both increase and display significant correlations before and after EVH and MCT, probably reflecting the same processes affecting temperature in the respiratory tract, independently of body temperature in general.

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The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

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