A Simple Device for Collecting Exhaled Breath Condensate (EBC) to Study Inflammatory Biomarkers of PM$_{10}$ Exposure in Thai Schoolchildren

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ABSTRACT

This study developed a portable device to collect exhaled breath condensate (EBC) and used it to collect EBC samples from schoolchildren exposed to ambient PM$_{10}$. The developed device was validated, including investigating the effect of collecting duration and breathing patterns on EBC volume, with five healthy volunteers. All five volunteers tolerated the device well, completing the EBC collection procedure without difficulty. Collecting normal tidal breathing for 10 minutes yielded the required EBC volume. We conducted a follow-up study with 104 healthy schoolchildren from two different primary schools in Chiang Mai, Thailand. We measured exhaled H$_2$O$_2$ concentrations in both the rainy and dry season; ambient PM$_{10}$ was significantly higher in the dry season. In the dry season, the mean exhaled H$_2$O$_2$ concentration was significantly higher in both groups (p<0.05). This study showed that the developed EBC collector device was cost effective, safe, rapid, and simple to use and exhaled H$_2$O$_2$ could be used as a biomarker for elevated PM$_{10}$ exposure before clinical symptoms appeared.

Keywords: Exhaled breath condensate collector device, Inflammatory marker, Schoolchildren, Chiang Mai, PM$_{10}$
INTRODUCTION

Previous studies have shown that airborne particulate matter less than 10 \( \mu \text{m} \) in aerodynamic diameter (PM\(_{10}\)) is a complex mixture of many pollutants – chemicals and transition metals, that are capable of redox cycling and that can stimulate inflammatory responses, especially in children (Kelly, 2003; Schwartz, 2004). Evidence has suggested that organic components deposited on the particle surface play an important role in mediating the toxic effect and inducing oxidative stress in the lungs, especially when antioxidant defenses have been overwhelmed. Airway inflammation plays an important role in the pathophysiology of various respiratory diseases. Respiratory tract health is traditionally assessed using airway biopsies, bronchoalveolar larvae (BAL) fluid, and bronchoscopy. Although these techniques provide direct information about the degree of airway inflammation, they are invasive, difficult, and not suitable for repeated use in children (Montushi and Barnes, 2003).

However, the precise mechanism of the relationship between ambient PM\(_{10}\) and respiratory health remains unclear, partly because of the lack of a non-invasive, biological sample, collection procedure for assessing lung inflammation. Recently, research has focused on analyzing biomarkers in exhaled breath. Exhaled breath condensate (EBC) is a new matrix for monitoring airway inflammation and oxidative stress markers of various respiratory conditions (Montushi and Barnes, 2003). EBC contains a number of volatile and non-volatile compounds derived from the respiratory surface, such as hydrogen peroxides, lipid peroxidation-derived products, and protein carbonyl groups (Taylor, 2011). In addition, EBC collection is non-invasive compared to bronchoscopy or induced sputum; it also facilitates repeated measurements in the same individual, a significant advantage over some of the other methodologies available for measuring inflammatory responses. Hydrogen peroxide (H\(_2\)O\(_2\)) is an oxidant produced by the alveolar membrane and can be measured in exhaled air. H\(_2\)O\(_2\), a marker of oxidative stress, is pathologically indicative of lung inflammation (van Beurden et al., 2002; Murata et al., 2014). Recently, numerous researchers have explored the utility of studying airway inflammation and oxidative stress in the context of pollution exposure (Jansen et al., 2005), as well as clinical monitoring (Antus and Kardos, 2015; Corradi et al., 2015; Garcia-de-la-Asuncion et al., 2015). EBC has been proposed as a simple, non-invasive tool for measuring airway inflammation. The biomarkers in EBC include H\(_2\)O\(_2\), which has been proposed as a method for assessing the health effects of air pollution in exposed populations (Doniec et al., 2005; Epton et al., 2008).

Although a variety of devices to collect EBC samples are commercially available, they are expensive and not suitable for field study. The developed EBC collecting device was based on the similar principle as the system of Dean et al. (2007), but with some modifications. The condensation chamber was constructed of polypropylene, following the recommendation of Horvath (2005) for EBC collection devices.

This study aimed to develop a portable device to collect EBC samples and, using these samples, assess airway inflammation in urban and highland school-children exposed to ambient PM\(_{10}\) in northern Thailand.
MATERIAL AND METHODS

Development and validation of a portable EBC collecting device

The portable EBC collecting device (Figure 1) consists of a mouthpiece with a one-way valve in which inspiratory and expiratory air are separated. The mouthpiece is connected to a flexible plastic tube (30 cm in length and 2 cm in internal diameter) that allows subjects to find a comfortable position. The tube connects to a 50 mL polypropylene collecting tube that acts as a sampling container. Polypropylene is a thermoplastic polymer used in variety of healthcare and laboratory applications, including syringes, tubing, hospital disposables, test tubes, beakers, and pipettes (Sastri, 2014). The collecting tube is placed inside a stainless steel chamber and designed to connect with a second one-way valve that allows the excess air in an expired breath to flow toward the top. A rubber ring between the flexible plastic tube and the hole in the stainless steel chamber creates an airtight connection. The stainless steel chamber contains liquid nitrogen in order to cool down the collecting tube.

![Diagram of the developed portable EBC collecting device.](image)

Figure 1. Schematic diagram of the developed portable EBC collecting device.

The developed device was validated for: (1) technical problems, such as any discomfort while wearing the device, (2) length of time required to collect adequate EBC volume, and (3) overall acceptance of the device by the volunteer subjects. Five healthy volunteers were recruited (12, 17, 25, 40, and 44 years old). Each volunteer was asked to breathe normally into the developed device for 10 minutes to collect EBC samples. After that, they were given a 30-minute break before collecting EBC samples a second time, for 20 minutes. Validation criteria
(1) and (3) were assessed by verbal communication between volunteer users and a trained research technician. For criteria (2), two EBC samples were collected and the condensate EBC volume of each sample was measured using a calibrated 1000-µL pipette.

**Application of the developed portable EBC device to collect EBC samples**

**Study site.** Two primary schools, one urban and one rural/highland, were selected as study sites. Chiang Mai Rajabhat University Demonstration School (or urban school) in Chiang Mai city represented the urban location. A recent study reported on the EBC malondialdehyde (MDA) as a biomarker of effect during elevated ambient PM10 levels at this study site (Phornwisetsirikun et al., 2014). Srinaeroo School (or highland school) represented the rural highland location.

The urban school is located in the northern part of Chiang Mai city, in the valley at about 300 m above mean sea level. This study site is adjacent to a street on one side and surrounded by workplaces and commercial areas on the other sides. The school is located within 2.5 km of the air quality monitoring station at the Yupparaj Wittayalai School in downtown Chiang Mai city. The PM$_{10}$ data from this station was assumed representative of the urban participants’ exposure to PM$_{10}$.

The highland school is located on Doi Suthep Mountain, about 35 km northwest of Chiang Mai city and about 1300 m above mean sea level. With no air quality monitoring station nearby, a portable airborne dust monitor (E-sampler, Met One Instruments Inc., USA) was used to collect PM$_{10}$ data at the location. This portable monitor was calibrated with the air quality monitoring station at the Yupparaj Wittayalai School, before using at the highland school. The PM$_{10}$ data from the portable airborne dust monitor was assumed representative of the highland participants’ exposure to PM$_{10}$.

**Criteria of study subjects.** To be included in the study, the schoolchildren had to have attended the selected primary school for at least one year and live within 2.5 km of the air quality monitoring station for the urban participants or the portable airborne dust monitor location for the highland participants. The children had to be 10-12 years old, not diagnosed with asthma or other chronic respiratory diseases, not on long-term medication, and willing to participate in the study.

The Human Experimentation Committee of the Research Institute for Health Sciences, Chiang Mai University, Thailand, approved the study protocol (Certificate HEC approval No. 1/2010). Children and parents signed written informed consent before participation.

**Study period.** The study was conducted during July 2011 (rainy season) and March 2012 (dry season).

**Exhaled H$_2$O$_2$ analysis**

EBC samples were collected from the schoolchildren using the developed device, as shown in Figure 2, during both the rainy season (low PM$_{10}$ level) and dry season (high PM$_{10}$ level). After rinsing their mouths, subjects were instructed
to form a complete seal around the mouthpiece and maintain a dry mouth during
collection by periodically swallowing excess saliva. The subjects sat comfortably
and wore nose clips. They were instructed to breathe normally and expire very
slowly through the mouthpiece; this continued for 10 minutes to collect conden-
sate of about 1.2 mL per subject. The collected EBC samples were immediately
stored at -70°C until analysis.

Figure 2. Collection of EBC samples using the developed device.

The concentration of $\text{H}_2\text{O}_2$ in EBC was measured using a spectrophotometric
assay by means of horseradish peroxidase-catalyzed oxidation of tetramethyl-
benzidine, according to the method previously described by Gallatin and Pratch
(1985). The detection limit was approximately 0.1 µM.

Data analysis
The statistical analysis was performed using the Statistical Package for
the Social Sciences for Windows (SPSS, Thailand) Version 17. The differences
between exhaled $\text{H}_2\text{O}_2$ levels and pulmonary function indices in the rainy and
dry seasons were determined using paired t-test. A p-value of less than 0.05 was
considered statistically significant.

RESULTS
Development and validation of a portable EBC collecting device
The developed EBC collecting device was validated with five healthy
volunteers; they reported no technical problems or complaints while wearing the
device. EBC samples from five healthy volunteers were collected and the mean
volumes by normally breathing over a period of 10 and 20 minutes were $1.24±0.07$
(n=5) and $1.30±0.05$ (n=5) mL, respectively (p=0.017). As 10 minutes yielded
adequate volume for the biomarker test in the present study, a collection time of
10 minutes was used throughout the study. The procedure of EBC collection using
the developed device was found safe, rapid, and simple to use and operate.
Application of the EBC device to collect EBC samples for investigating the airway inflammation of schoolchildren

General characteristics of the study participants. Table 1 shows the general characteristics of the participants. The participants, all between the ages of 10-12 years, had a median height of 129-144 cm and weight of 27-34 kg. However, the urban children were significantly taller and heavier than their highland counterparts, partly skewed by one overweight subject of 59 kg.

Table 1. General characteristics of the participants.

<table>
<thead>
<tr>
<th>Participant characteristics</th>
<th>Urban school (n = 54)</th>
<th>Highland school (n = 50)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male : Female</td>
<td>27 : 27</td>
<td>33 : 17</td>
<td>0.101</td>
</tr>
<tr>
<td>Age, year **</td>
<td>11 (10-12)</td>
<td>11 (10-12)</td>
<td>0.213</td>
</tr>
<tr>
<td>Height, cm **</td>
<td>144 (30-164)</td>
<td>129 (113-155)</td>
<td>0.000*</td>
</tr>
<tr>
<td>Weight, kg **</td>
<td>34 (25-59)</td>
<td>27 (20-50)</td>
<td>0.000*</td>
</tr>
</tbody>
</table>

Note: *significant difference between schools (p < 0.05), ** median (range).

PM$_{10}$ and exhaled H$_2$O$_2$ concentrations. In both schools, the 24-hour mean PM$_{10}$ in the dry season was significantly higher than in the rainy season. However, it was five times higher at the urban school, while only twice as high at the highland school; this difference between these two locations was also statistically significant. Both levels of PM$_{10}$ during the study period did not exceed Thailand’s 24-hour mean PM$_{10}$ limit of 120 µg/m$^3$ (Thailand Air Quality and Noise Standards, 2004). This phenomenon has been explained by the westerly wind that blows into Chiang Mai City (Wiriy et al., 2013). Also, Doi Suthep Mountain, to the west, is a National Park and open biomass burning is strictly prohibited.

At the urban school, the mean concentration of exhaled H$_2$O$_2$ was 0.17 µM in the rainy season and 0.21 µM in the dry season (p = 0.003). At the highland school, the mean concentration of exhaled H$_2$O$_2$ was 0.16 µM in the rainy season and 0.18 µM in the dry season (p=0.001). The concentrations of exhaled H$_2$O$_2$ in both schools increased significantly in the dry season.

Table 2. Comparison of PM$_{10}$ and exhaled H$_2$O$_2$ concentrations of the schoolchildren from urban and highland schools between the rainy and dry seasons.

<table>
<thead>
<tr>
<th>Variables</th>
<th>Mean ± SD</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rainy season</td>
<td>Dry season</td>
</tr>
<tr>
<td>Urban School</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean 24 h PM$_{10}$ (n=5), µg/m$^3$</td>
<td>16.7 ± 1.3</td>
<td>90.7 ± 27.0</td>
</tr>
<tr>
<td>Exhaled H$_2$O$_2$ conc. (n=54), µM</td>
<td>0.17 ± 0.08</td>
<td>0.21 ± 0.08</td>
</tr>
<tr>
<td>Highland School</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean 24 h PM$_{10}$ (n=5), µg/m$^3$</td>
<td>22.6 ± 1.0</td>
<td>50.6 ± 8.8</td>
</tr>
<tr>
<td>Exhaled H$_2$O$_2$ conc. (n=50), µM</td>
<td>0.16 ± 0.04</td>
<td>0.18 ± 0.05</td>
</tr>
</tbody>
</table>

Note: *significant difference between schools (p < 0.05).
DISCUSSION

We developed a cost-effective EBC collecting device for in-house and field use; each device costs about THB 600 (about USD 20) (Figure 2) and consumes liquid nitrogen for cooling of about THB 5 per sample. The device was employed in a field study of 104 primary schoolchildren in 2011-2012 in Chiang Mai. The collection procedure, which took only 20 minutes per person, was simple and did not require skilled medical staff. A trained operator was sufficient to collect the EBC samples from the subjects. Our collection procedure compared favorably with Mutlu et al. (2001), who reported that collection usually takes 5-10 minutes in adults and up to 15-20 minutes in children to obtain 1-3 mL of EBC. Furthermore, our developed device, which can be used in the field, offered similar performance to commercially available devices used in research and clinical studies, including children with respiratory diseases, which collect 1.5-2 mL of EBC in 10-15 minutes (Romieu et al., 2008; De Prins et al., 2014; Rosa et al., 2014).

Exhaled $H_2O_2$, the inflammatory biomarker used in our study, has been shown to be a significant biomarker from ambient elevated PM$_{10}$ exposure. Our results here also confirm our previous findings, which reported that the EBC malondialdehyde (MDA) biomarker of oxidative stress was raised in children exposed to PM$_{10}$ air pollution (Phornwisetsirikun et al., 2014). These results are also consistent with other studies of EBC biomarkers of pulmonary inflammation (Ralph et al., 2006; Gergelova et al., 2008; Chow et al., 2009).

Although possible confounding factors may exist in measuring exhaled $H_2O_2$ concentration, the present study design was a follow-up study of the same person, in order to minimize individual variability. Therefore, the present study results demonstrated that the exhaled $H_2O_2$ concentrations, as well as the MDA concentrations from our previous report (Phornwisetsirikun et al., 2014), are suitable biomarkers of elevated PM$_{10}$ exposure. In conclusion, our study developed an economical device for collecting EBC that was non-invasive, safe, rapid, and simple to use. In addition, the results indicated that exhaled $H_2O_2$ concentration provided good information about inflammation in the respiratory system of children (i.e., healthy schoolchildren in the present study) before clinical symptoms appear.

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