Research Article

Influence of Milk on Exhaled Carbon Monoxide (CO) Measurement by Portable CO Monitors

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Background. A portable breath carbon monoxide (CO) monitor has a high cross-sensitivity to hydrogen (H2). This study examined the influence of H2 after consuming milk on the detected CO values using three types of portable CO monitors.

Materials and Methods. Exhaled breath from seven participants (four healthy nonsmokers and three smokers with otherwise unknown comorbidities) was collected in sampling bags. The participants then consumed 200 mL of milk, and the exhaled breath of each was collected in separate bags every 30 minutes until 9 hours later. CO and H2 in the bag were measured using a gas chromatograph as a reference analyzer, and CO was also measured using three types of portable CO monitors.

Results. After consuming milk, H2 levels were significantly higher, and CO levels were not significantly elevated as measured by the reference analyzer. However, CO levels in monitors A and B were significantly elevated, even though participants did not smoke. The H2 levels in the reference analyzer significantly increased and reached a maximum 4.5 hours after consuming milk. The difference in CO levels between the reference analyzer and each monitor increased significantly after 5 or 5.5 hours.

Conclusions. This study suggested that the breath CO monitors with a cross-sensitivity to H2 responded to H2 as CO in the exhaled gas and measured higher than actual values after milk consumption. The extent of the influence of H2 differed depending on the type of CO monitor. It is necessary to consider milk consumption when assessing the smoking status of people using portable CO monitors.

1. Introduction

One of the toxic substances produced by smoking is carbon monoxide (CO), a product of converting heme to biliverdin by heme oxygenase in microsomes. CO binds to heme molecules such as hemoglobin, causing tissue hypoxia and oxidative stress [1].

An easy and objective method for evaluating smoking status is to measure the CO in smokers’ breath using a CO monitor, which is the “stethoscope” of a tobacco treatment specialist. It is also used to measure the nicotine dependence level and passive smoke exposure and plays an important role in the titration of combination medication dosing. It is a powerful motivational tool for quitting smoking. Currently, three types of portable breath CO monitors are used to assess smoking status in smoking cessation programs [2–4]. It has been reported that the sensor of the CO monitors has a high cross-sensitivity to hydrogen (H2). In people with hypolactasia or lactose intolerance, lactose in milk, which is not metabolized in the small intestine, is changed into H2 by the intestinal bacteria and is absorbed. In such cases, the portable CO monitors may erroneously measure H2 as CO in the exhaled gas and measured higher than actual values after milk consumption. The extent of the influence of H2 differed depending on the type of CO monitor. It is necessary to consider milk consumption when assessing the smoking status of people using portable CO monitors.
has gastrointestinal symptoms [8], and H$_2$ is often detected in the breath of people who consume milk or lactose but do not have such symptoms.

In this study, to properly assess CO levels detected in the portable breath CO monitor, we examined the influence of consuming milk on the CO values using three types of monitors and compared the values with that of a reference analyzer.

2. Materials and Methods

2.1. Participants. Participants were seven individuals (four healthy nonsmokers and three smokers with otherwise unknown comorbidities) without respiratory diseases and milk allergies. Moreover, they were not diagnosed as lactose intolerant. They were either students or affiliated with a Japanese university and recruited in a class on health as volunteers between December 2019 and December 2020. Participants with obvious milk allergies and lung or bronchial abnormalities were excluded. Written informed consent was obtained from all participants.

2.2. Procedure. Participants consumed no milk and dairy products the day before the study. After an overnight fast (only water intake was allowed) and nonsmoking, at 8:30 a.m., participants exhaled and inhaled completely, held their breath for 15 seconds, and then exhaled rapidly into a 600 mL sampling bag (Taiyo Corporation, Osaka, Japan) in which the exhaled air was collected. At 9 a.m., participants consumed 200 mL of milk (containing 8.6 g lactose). After consuming milk, the exhaled breath of every participant was collected in a sampling bag every 30 minutes until 5 p.m., in the same manner. Participants did not consume any food or drink and were prohibited from smoking during the study. The participants were asked about their subjective gastrointestinal symptoms to evaluate them as lactose intolerant after consuming milk. The study design is shown in Figure 1.

A gas chromatograph with a semiconductor detector (TRIlyzer mBA-3000, Taiyo Corporation, Osaka, Japan) was used as a reference analyzer. Some of the exhaled breath was injected into the reference analyzer from each sampling bag, and CO and H$_2$ were measured. The remaining exhaled air in the bag was injected into three different types of portable CO monitors (monitor A: Smokerlyzer (PICOplus®), Bedfont Inc.; monitor B: Micro CO monitor, Vyaire Medical Inc.; and monitor C: Smokerlyzer (PICO advance®)). Before the study, the analyzer was calibrated with a mixture of CO and air [9]. CO values were measured by the reference analyzer and three monitors, and H$_2$ values were measured by the reference analyzer. CO and H$_2$ values were compared in time series.

2.3. Statistical Analysis. Data were analyzed using IBM SPSS version 25 (IBM Corp., NY, USA). The Wilcoxon signed-rank test was used to compare the paired groups (levels before consuming milk and at each time after consuming milk). Spearman’s correlation coefficient was used to evaluate the relationship between the two parameters. Data are presented as medians (first and third quartiles). Statistical significance was set at $P < 0.05$. Significant differences were declared at $P \leq 0.05$ and tendencies between $0.05 < P \leq 0.1$.

The study was approved by the Research Ethics Committee of Kyoto Women’s University (approval number 2019-25) and was conducted in accordance with the guidelines of the Declaration of Helsinki.

3. Results

Table 1 shows the characteristics of the participants and CO and H$_2$ levels before and after consuming milk. Participants were aged between 22 and 60 years, five men and two women, three smokers, and four nonsmokers. Only one had occasional abdominal symptoms after consuming milk; however, the participant had no symptoms in this study.

Before consuming milk, participants’ CO levels ranged from 1.0 to 16.4 ppm by the reference analyzer and from 1 to 9 ppm, 0 to 13 ppm, and 1 to 10 ppm in monitors A, B, and C, respectively. H$_2$ levels in the reference analyzers ranged from 1.6 to 19.1 ppm. Participants had no abdominal symptoms after consuming milk in this study. After consuming milk, the CO levels of almost all of the participants increased compared with those before consuming milk. H$_2$ levels increased 1.8- to 16.8-fold compared to those before consuming milk.

Table 2 shows CO levels measured by monitors A, B, and C and the reference analyzer and H$_2$ levels measured by the reference analyzer before and after consuming milk. No significant differences were found in the median CO levels measured by the reference analyzer and monitors A, B, and C before milk consumption. After milk consumption, H$_2$ levels in the reference analyzer were significantly higher.
<table>
<thead>
<tr>
<th></th>
<th>Sex</th>
<th>Age</th>
<th>Smoker or not</th>
<th>Habitual symptoms of lactose intolerance</th>
<th>Symptoms during the study</th>
<th>CO levels (ppm) before consuming milk</th>
<th>Reference H₂ levels (ppm)</th>
<th>Maximum CO levels (ppm) after consuming milk</th>
<th>Maximum H₂ levels after consuming milk</th>
<th>Multiples compared to reference H₂ levels</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>M</td>
<td>23</td>
<td>No</td>
<td>Sometimes</td>
<td>Nothing</td>
<td>1.0</td>
<td>3.8</td>
<td>3.0</td>
<td>1.6</td>
<td>28.4</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>60</td>
<td>No</td>
<td>Nothing</td>
<td>Nothing</td>
<td>3.0</td>
<td>2.0</td>
<td>4.0</td>
<td>2.0</td>
<td>16.83</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>22</td>
<td>No</td>
<td>Nothing</td>
<td>Nothing</td>
<td>2.0</td>
<td>1.0</td>
<td>2.0</td>
<td>1.0</td>
<td>15.0</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>22</td>
<td>No</td>
<td>Nothing</td>
<td>Nothing</td>
<td>2.0</td>
<td>3.0</td>
<td>2.0</td>
<td>1.0</td>
<td>35.3</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>27</td>
<td>Smoker</td>
<td>Nothing</td>
<td>Nothing</td>
<td>7.0</td>
<td>2.0</td>
<td>4.0</td>
<td>2.0</td>
<td>39.7</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>25</td>
<td>Smoker</td>
<td>Nothing</td>
<td>Nothing</td>
<td>4.0</td>
<td>6.0</td>
<td>6.0</td>
<td>4.0</td>
<td>21.7</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>25</td>
<td>Smoker</td>
<td>Nothing</td>
<td>Nothing</td>
<td>9.0</td>
<td>13.0</td>
<td>12.0</td>
<td>12.0</td>
<td>36.6</td>
</tr>
</tbody>
</table>
Table 2: CO levels measured by monitors A, B, and C and the reference analyzer and H₂ levels measured by the reference analyzer before and after consuming milk.

<table>
<thead>
<tr>
<th></th>
<th>Before consuming milk</th>
<th>Maximum levels after consuming milk</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO levels (ppm)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Monitor A</td>
<td>3.0 (2.0, 7.0)</td>
<td>4.0 (2.0, 8.0)</td>
<td>0.039*</td>
</tr>
<tr>
<td>Monitor B</td>
<td>3.0 (1.0, 7.0)</td>
<td>5.0 (4.0, 9.0)</td>
<td>0.026*</td>
</tr>
<tr>
<td>Monitor C</td>
<td>2.0 (1.0, 7.0)</td>
<td>2.0 (2.0, 8.0)</td>
<td>0.059</td>
</tr>
<tr>
<td>Reference</td>
<td>2.5 (1.2, 7.7)</td>
<td>2.6 (1.2, 9.8)</td>
<td>0.058</td>
</tr>
<tr>
<td>Reference H₂ levels (ppm)</td>
<td>6.2 (3.8, 17.4)</td>
<td>28.4 (21.7, 36.6)</td>
<td>0.018*</td>
</tr>
</tbody>
</table>

Values are expressed in median (first quartile and third quartile). *P < 0.05.

Figure 2: Time course of expiratory H₂ levels. Expiratory H₂ levels gradually increase after consuming milk and are significantly higher than those before consuming milk. *P < 0.05 vs. the levels before consuming milk.

(P = 0.018) and CO levels in the reference analyzer were not significantly elevated. However, CO levels in monitors A and B were significantly elevated (P = 0.039 and P = 0.026, respectively).

Figure 2 shows the time course of H₂ levels in the reference analyzer. The H₂ levels in the reference analyzer significantly (P = 0.018) increased after consuming milk and reached a maximum at 13:30 (4.5 hours after milk intake).

Figure 3 shows the relationship between the difference in CO levels of each monitor and the reference analyzer and the expiratory H₂ levels. The difference between the CO values of each monitor and the reference analyzer was significantly correlated with the H₂ levels of the reference analyzer. The difference in monitor B had the strongest correlation with the H₂ levels of the reference analyzer. In monitor B, approximately one-tenth of the hydrogen concentration in the exhaled air was mistakenly measured as CO levels.

Figure 4 shows the time course of the difference in CO concentration between each monitor and the reference analyzer. The difference in CO levels between the reference analyzer and each monitor showed an increasing trend after consuming milk in all monitors. In monitors A and B, it became significant at 14:00 (5 hours later). In monitor C, it became significant at 14:30 (5.5 hours later).

4. Discussion

This study illustrated that after consuming 200 mL of milk, the CO levels detected in monitors A and B increased significantly after 5 to 5.5 hours compared to the values before consumption, despite the fact that participants did not smoke. Individual differences were observed in the levels and duration of the increase in CO levels measured by the monitors. This is the first study to show the effect of H₂ produced by consuming milk on the values measured by different models of portable CO monitors.

All three portable CO monitors used in this study use electrochemical gas sensors. The electrochemical analysis method measures the electric current produced in an aqueous solution by electrical oxidation by an electrode that has acted as a catalyst [10, 11]. As detailed measurement methods are not disclosed by the manufacturer, there was no consistent agreement on the CO value between models [3]. As this electrochemical sensor also reacts with H₂, hydrogen sulfide, sulfur dioxide, nitrogen dioxide, nitrogen monoxide, and ethylene, if there is H₂ in the expired breath, it may be erroneously measured as CO [5]. The instructions for the Bedfont Scientific Ltd. instrument describe the possibility of H₂ crossover (interference with H₂) but do not describe the specific effects of lactose ingestion (time and extent).

The activity of lactase, a lactose-degrading enzyme at the brush border of the small intestinal mucosal epithelium, is deficient or reduced in people with hypolactasia or lactose intolerance. Therefore, lactose, a disaccharide, is not degraded into glucose and galactose. The lactose that cannot be degraded is not absorbed in the small intestine and is fermented by intestinal bacteria in the large intestine, and H₂ is produced. The produced H₂ is absorbed through the intestinal mucosa, dissolved into the blood, and diffused into the alveoli via the blood circulation, and some of it is expired in the exhaled air [12]. Lactose also irritates the large intestine, causing the abdominal symptoms of lactose intolerance. The objective evaluation of lactose intolerance is done by measuring enzyme (lactase) activity using biopsy material of the small intestine [13]. In the participants of this study, a possibility of low lactase activity (subclinical lactose intolerance [14]) was suggested because of the H₂ in their exhaled breath after consuming milk, although they were not aware of lactose...
intolerance. Therefore, before measuring CO levels by the breath CO monitor, it is necessary to check the consumption of milk or foods that may produce H₂, the time after consumption, and the type of monitor used.

After consuming milk, the increase in CO levels detected in monitor C was less than in A and B. This is because the portable monitor C is the most recent model; hence, the influence of H₂ may be minimal due to advances in technology including calibration adjustment of the monitor. However, CO monitor C might have measured a value slightly lower than the actual value.

Few studies have examined the effect of H₂ on CO monitor readings. We have reported in a previous study that in eleven nonsmokers who consumed 400 mL of milk, the levels of the three portable CO monitors were significantly elevated from 1.5 hours to a maximum of 8 hours after ingestion, up to a maximum of 18 ppm [8]. Our previous report showed a greater degree of elevation of the CO levels compared to the present study. This is presumably because 200 mL of milk was consumed in the present study, whereas 400 mL of milk was consumed in the previous one, suggesting a dose dependence on the relationship between the amount of H₂ produced and amount of lactose ingested. Another previous report using the Bedfont Micro Smokerlyzer monitor on four lactose-intolerant persons showed that an H₂ concentration of 38.91 ppm in exhaled air was sufficient to record a CO level of 10 ppm, and this level is equivalent to the ingestion of 350 mL of milk [5].

There are several limitations to this study. First, various foods other than lactose produce H₂ in the intestine, but this study focused only on lactose. Even when foods that do not require digestive enzymes, such as dietary fiber and indigestible carbohydrates, are ingested, they may pass undigested through the small intestine and be fermented by intestinal bacteria in the large intestine, producing H₂ [15, 16]. Some participants in this study had elevated H₂ levels before milk consumption, which might be related to the previous day’s diet. Second, in the study, exhaled air was injected into the sampling bag and measured by a reference analyzer and CO monitor. Therefore, there may be a difference in levels.
compared to those obtained when exhaled air is directly blown into the CO monitor. Third, this study only examined a time course of up to 8 hours. Fourth, we have not been able to examine in detail the causes of H2 production, such as the degree of decrease in lactase activity and differences in the state of the intestinal bacteria. Fifth, the sample size of this study is small; therefore, careful consideration should be given to realistic application in clinical practice. The causes of individual differences could not be examined because of the small number of participants. Sixth, the relation between the nicotine dependence level of the smokers and their expired CO levels was not investigated. Seventh, the study lacked a control group who did not drink milk.

5. Conclusions

The results of this study showed that when a portable CO monitor was used to measure CO after lactose intake, the CO monitor responded to H2, and the measured value increased even if the exhaled air did not contain CO, regardless of whether the participant had subjective symptoms of lactose intolerance or not. The extent of the effect differed depending on the type of CO monitor. Therefore, when assessing the smoking status using portable breath CO monitors, it is necessary to consider prior consumption of milk or foods that may produce H2, the time after consumption, and the type of monitor used. Further studies are needed to explore the influences of foods that may produce H2 on the CO values using portable CO monitors.

Data Availability

Data are available from the corresponding author on reasonable request.

Conflicts of Interest

The authors declare that there is no conflict of interest regarding the publication of this article.

Acknowledgments

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References