Factors influencing breath condensate volume

Abstract

Analysis of breath condensate (BC) has received interest recently due to the need for easy and repetitive monitoring of airway and pulmonary disease. While many authors have used custom built systems, commercial systems are now available and will probably be used more widely. Early studies of markers and mediators in BC have reported concentrations following varying periods of sampling time. However, factors that influence the generation of BC have not been analysed and it is unclear whether breathing rate, tidal volume, lung function, body weight, height or age influence the amount of BC collected. We therefore studied the influence of these factors on breath condensate volume and breath condensate urea and protein concentrations in 22 healthy volunteers and 23 COPD patients.

A strong correlation of total respired volume and breath condensate volume was observed for both groups (volunteers: $r = 0.952$, $p < 0.0001$, COPD: $r = 0.883$, $p < 0.0001$) while no significant correlation existed for breath condensate volume and TLC, RV, VC, FEV1, $R_{exh}$, height or body weight. As long as ventilation remained fairly constant, breath condensate volume increased linearly with time. The fraction of breath condensate extracted from total vapour contained in the exhale was estimated by measuring relative atmospheric humidity before and after the collecting tube. The amount calculated by the change in temperature and saturation corresponded closely to the amount actually collected.

We conclude from these results that breath condensate volume is primarily dependent on $V_{E}$, and does not seem to depend on lung function parameters. For standardisation it is suggested to report breath condensate measurements per volume respired. Both, urea and protein are present in measurable quantities in breath condensate and protein as well as BCV may be helpful denominators for comparison with e.g. cytokines in lung disease.

Introduction

Breath condensate has been suggested to be a non-invasive method of sampling the epithelial lining fluid (ELF) or more precisely that part of the ELF that finds its way into the exhalate. Regardless whether it truly reflects ELF, breath condensate contains not only soluble small molecules and hydrogen peroxide [3], but also a number of large molecules such as leukotrienes B4, C4, D4, E4, Prostaglandin E2 ECP, IL-1 siL-2R, TNF-, isoprostane, tumour markers and small amounts of DNA [1,4–7,15]. BC evaluation may be helpful in determining the degree of airway inflammation or epithelial damage, investigating surfactant integrity, tumour growth and possibly genome mutations [4,5]. Breath condensate has been used for analysis of leukotrienes in inflammatory upper airway disease [8], $H_{2}O_{2}$ and pH in asthma [2,9], several markers of inflammation in COPD [10,11], and mechanically ventilated patients, patients with acute respiratory failure [12,13] and cigarette smokers [14]. In addition to the variety of parameters that may be monitored in breath condensate it’s attractiveness is the ease of sample collection. This allows frequent follow up and improved control of lung disease when appropriate markers have been identified. In addition breath condensate monitoring is less expensive than more invasive ways of monitoring lung disease.

Most likely, breath condensate is a mixture of water vapour and aerosolised ELF, but the exact origin and mechanism of breath condensate generation remains to be determined. Collection of this material involves rapid cooling of the exhale with condensation of water vapour as well as sedimentation and attachment of aerosol particles to very cold surfaces. The relative importance of extraction by cooling (condensate formation) and sediment due to gravity or impact for the process of condensate formation has been debated. Complete extraction from the exhale to a freezing cold surface will depend on the size and shape of the velocity of the airstream passing the cold surface and the temperature gradient [8].

For the future it will be useful to define factors influencing breath condensate formation and ways for standardization of breath condensate collection. Without sufficient knowledge of influent parameters, breath condensate collection will suffer from many of the drawbacks of examining solutes in bronchoalveolar lavage fluid [16].

We here investigated the influence of ventilation ($V_{E}$) and lung function as well as individual parameters on breath condensate volume. In addition we examined the effectiveness of breath condensate collection by determining humidity and temperature in the exhale before and after the cold trap.
Material and methods

Breath condensate was collected using the ECoSreen® (Jaeger/Toennies, Wuerzburg, Germany). This device has a cold trap with one collecting tube (Ct-1) inserted into a receptacle integrated into the cooled head sampling unit. A second cold trap was fitted to the exhalation port of the first one. This second trap called collecting tube II (Ct-2) was cooled to ~20 °C by immersion into dry ice in ethanol. The outlet of this second cold trap was open to the surrounding (Fig. 1).

Breath condensate was collected during three consecutive 10 min periods. “Normal breathing” was requested by the unit operator, but intentionally no influence on rate or tidal volume was exerted. For each of the 10 min periods, aliquots were stored separately after briefly thawing and collecting the entire fluid into the detachable part of the collecting tube. For each of the three collection periods the tube in Ct-1 position was exchanged, while in the Ct-2 position only one collection tube was used for the entire time sampling period of 30 min. In total four different aliquots were therefore collected from each individual.

Measurement of $V_e$ was performed with a bellows type gas flow counter (Model BK4, Krom Schröder, Germany; $Q_{min}$: 0.04 L/min, $Q_{max}$: 6.0 L/min; accuracy ±1.5 % at 20 °C) fitted to the inspiratory opening. 22 healthy volunteers and 23 patients with a COPD underwent a complete lung function study.

Humidity was measured using a precision hygrometer (P 570; Novodirect, Germany; sensor precision 1 % for humidity, 1 °C for temperature, sensor resolution 0.1 % for humidity, 0.1 °C for temperature) with a minimum registration interval of 1 s. Smartgraph 1.07 and De Terminal P 500 software was used for evaluation (Dostmann electronic GmbH, Wertheim, Germany).

Relative humidity and temperature were measured at the inlet and outlet of the collecting tube consecutively. $V_e$ was calculated from $V_i$ measured at the inspiratory port using the equation: $V_i/T_i = V_e/T_e$. The amount of water in the air entering and leaving the collecting tube was calculated [17] and plotted versus time (100 s; n = 8 volunteers; example in Fig. 2). The area under the resulting curve (AUC) thus represented the amount of water that had entered/extracted the collecting tube. The calculated difference represented the extracted fraction. The fraction of water was then compared to the amount actually collected. The difference between these two was interpreted to be due to incomplete yield from the cold trap after the process of thawing and pipetting.

Protein concentration contained in breath condensate was measured using the Micro BCA Protein Assay (Pierce, Rockford USA) with a linear working range of 0.5 – 20 μg/ml [18,19]. For the reason of comparison, samples from Ct-1 and Ct-2 were analysed separately.

In order to exclude a possible contamination of breath condensate by saliva we measured amylase activity in breath condensate of all volunteers and patients. As a reference value for amylase activity in saliva we used the average saliva value of five volunteers. Amylase activity was determined with the alpha-Amylase ESP 1491300 kit (test limit 0.03 μmol/l's; Boehringer Mannheim, Germany). For quantitative determination of urea nitrogen in breath condensate we used a BUN test kit from Sigma (Deisenhofen, Germany).

Statistical analysis was performed with the SPSS software package (SPSS Inc., Chicago, USA). Linear regression analysis was used for analysing the influence of $V_e$ and lung function variables on breath condensate volume from each of the three sampling periods. Inter-individual as well as intra-individual comparisons of condensate volumes from all the sampling periods were performed by ANOVA testing. p was considered significant at the 5 % level. Results are mean ± SD.

Results

Breath condensate volumes and ventilation of volunteers and COPD patients are listed in Table 1. $V_e$ ranged from 37 – 302 L in the volunteer group and from 27 – 238 L in the COPD group (all 10 min periods included). We observed no significant variation of $V_e$ within individuals comparing the first, second
Table 1  BCV and \( V_v \) in COPD and volunteers

<table>
<thead>
<tr>
<th>Sampling period</th>
<th>BCV volunteers</th>
<th>COPD</th>
<th>( p )</th>
<th>( V_v ) volunteers</th>
<th>COPD</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 – 10 min</td>
<td>0.94 ml ± 0.53</td>
<td>1.22 ml ± 0.57</td>
<td>0.10</td>
<td>102.6 L ± 53.3</td>
<td>129.5 L ± 62.1</td>
<td>0.13</td>
</tr>
<tr>
<td>11 – 20 min</td>
<td>0.93 ml ± 0.43</td>
<td>0.98 ml ± 0.54</td>
<td>0.71</td>
<td>96.4 L ± 43.4</td>
<td>105.4 L ± 58.2</td>
<td>0.58</td>
</tr>
<tr>
<td>21 – 30 min</td>
<td>0.97 ml ± 0.54</td>
<td>1.13 ml ± 0.47</td>
<td>0.49</td>
<td>102.4 L ± 59.3</td>
<td>120.7 L ± 59.6</td>
<td>0.48</td>
</tr>
<tr>
<td>average</td>
<td>0.94 ml ± 0.49</td>
<td>1.12 ml ± 0.55</td>
<td>0.09</td>
<td>100.5 L ± 51.8</td>
<td>118.9 L ± 60.1</td>
<td>0.08</td>
</tr>
</tbody>
</table>

Date shown as mean ± SD

and third 10 min periods (data not shown) and there was also no significant within-subject variability of BCV (mean ± SD in volunteers: 0.95 ml ± 0.14, \( p = 0.96 \); mean ± SD in COPD patients: 1.18 ml ± 0.15, \( p = 0.38 \)).

The correlation of BCV and \( V_v \) in the first cold trap (Ct-1) for the entire 30 min sampling period for volunteers and COPD patients is shown in Fig. 3. Correlation of BCV and \( V_v \) for the second cold trap (Ct-2) was also very strong for volunteers (\( y = 75.3x + 44.4, r = 0.95, p < 0.0001 \)) as well as for COPD patients (\( y = 91.0x + 72.9, r = 0.87, p < 0.0001 \)). Comparably strong correlations were observed when each of the 10 min periods was analysed separately (volunteers: first 10 min period: \( r = 0.91, p < 0.0001 \), second 10 min period: \( r = 0.90, p < 0.0001 \), third 10 min period: \( r = 0.92, p < 0.0001 \); COPD patients: 0 – 10 min: \( r = 0.75, p < 0.0001 \), 10 – 20 min: \( r = 0.83, p < 0.0001 \), 20 – 30 min: \( r = 0.94, p < 0.001 \)). The increase of BCV as well as the increase of \( V_v \) with time was parallel. Table 1 shows BCV and \( V_v \) in volunteers and COPD patients. The ratio of \( V_v/BCV \) was almost identical in both groups (data not shown). COPD patients exhibited an increased, compensatory ventilation leading to a concomitant increase in BCV.

A \( V_v \) of 100 L resulted in a BCV of 0.95 ml ± 0.24 in volunteers and 1.02 ml ± 0.39 in COPD patients. The relation of \( V_v/BCV \) did not vary significantly in the three sampling periods.

None of the lung function parameters measured (TLC, VC, RV, FEV1 and \( R_{et} \)) correlated significantly with BCV in either volunteers or COPD patients (Table 2). Age and bodyweight were not significantly influential on breath condensate volume (Table 2).

The efficiency of breath condensate collection was evaluated in eight consecutive volunteers. In these individuals mean temperature and relative humidity at the inlet of Ct-1 was 29°C ± 0.41 and 88.05% ± 4.44 respectively. The corresponding mean values at the outlet of Ct-1 were 19°C ± 0.5 and 83.99% ± 3.5 (Fig. 4).

The percentage of water extracted from breath condensate by Ct-1 was 36.9% ± 6.1 in volunteers and 40.7% ± 14.9 in COPD patients. Comparably the second cold trap in a row extracted 43.7% ± 7.8 in volunteers and 30.2% ± 11.5 in COPD patients. All together with both traps 80.6% ± 10.5 was extracted in volunteers and 70.9% ± 17.2 in COPD patients. Thus no difference existed in the efficiency of water vapour extraction in healthy individuals versus COPD patients.

Mean protein concentrations in all breath condensate aliquots were 11.9 µg/ml ± 8.9 in volunteers and 12.9 µg/ml ± 7.5 in COPD patients. Concentrations in aliquots of Ct-1 were similar to those of Ct-2 (volunteers: Ct-1 10.6 µg/ml ± 6.2 vs. Ct-2 13.3 µg/ml ± 10.9; \( p = 0.32 \); COPD patients: Ct-1 13.5 µg/ml ± 9.3 vs. Ct-2 12.4 µg/ml ± 5.5; \( p = 0.62 \)). Total protein correlated significantly with BCV (volunteers: \( r = 0.78, p < 0.0001 \); COPD patients: \( r = 0.46, p = 0.027 \) as well as \( V_v \) (volunteers: \( r = 0.65, p < 0.001 \); COPD patients: \( r = 0.49, p = 0.017 \); Fig. 5A). Because BCV and \( V_v \) both were strongly correlated with protein concentration, our findings suggest that protein accumulated by a similar mechanism as breath condensate. The weaker correlation in COPD patients may indicate additional or reduced mechanisms of protein entering the cold trap in COPD patients.
Factors influencing breath condensate volume

<table>
<thead>
<tr>
<th>parameters</th>
<th>volunteers (collecting unit 1)</th>
<th>COPD patients (collecting unit 1)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R</td>
<td>p</td>
</tr>
<tr>
<td>age [year]</td>
<td>0.31</td>
<td>0.16</td>
</tr>
<tr>
<td>bodyweight [kg]</td>
<td>0.37</td>
<td>0.09</td>
</tr>
<tr>
<td>height [cm]</td>
<td>0.20</td>
<td>0.37</td>
</tr>
<tr>
<td>TLC*</td>
<td>0.28</td>
<td>0.20</td>
</tr>
<tr>
<td>VC**</td>
<td>0.24</td>
<td>0.29</td>
</tr>
<tr>
<td>RV***</td>
<td>0.32</td>
<td>0.14</td>
</tr>
<tr>
<td>FEV1##</td>
<td>0.04</td>
<td>0.87</td>
</tr>
<tr>
<td>R tot#</td>
<td>0.18</td>
<td>0.41</td>
</tr>
<tr>
<td>volume expired [L]</td>
<td>0.96</td>
<td>0.0001</td>
</tr>
</tbody>
</table>

* total lung capacity [L]; ** vital capacity [L]; *** residual volume [L]; ## forced expiratory volume [L]; # airway resistance [kPa·s/L]

**Table 2** Correlation of biological and lung function parameters with breath condensate volume.

Activity of alpha-amylase was undetectable in BCV of volunteers or COPD patients. The reference value of alpha-amylase activity in saliva (measured in five volunteers) was 7.3 µmol/l·s ± 4.9.

Measurement of urea concentration was performed in breath condensate from Ct-1 as well as Ct-2. Mean urea concentration was significantly elevated in COPD patients (370.4 µmol/l ± 244.21) compared to healthy volunteers (83.6 µmol/l ± 56.3). Correlating the absolute amount of urea with BCV and V E we observed a linear regression for urea and BCV (r = 0.91; p < 0.0001) as well as for urea and V E (r = 0.92; p < 0.0001; Fig. 5B) in volunteers only.

**Discussion**

In this study of 22 healthy volunteers and 23 COPD patients we have demonstrated that breath condensate volume depends primarily on ventilation. Lung function, body weight, height, and age did not correlate significantly with the amount of breath condensate recovered. A correlation of BCV with ventilation might have been expected, however we did not foresee the lack of other influential factors in these two populations of healthy individuals and COPD patients. In addition, the correlation of BCV and ventilation is extraordinarily strong and leaves little room for other influential factors.

Although it may seem of minor importance that BCV is predominantly dependent on ventilation, this finding will facilitate breath condensate evaluation to a great extent. Inter- or even intra-individual variability in the amount of breath condensate volume in relation to ventilation would have suggested different mechanisms of breath condensate generation with varying relative importance. For example the role of aerosol admixture to aqueous vapour could have been greatly variable. This in turn would have suggested a varying composition of the condensate and a need for a close control of ventilation parameters. However with a linear and strong correlation of breath condensate volume in healthy volunteers with ventilation over a rather wide range it is unlikely that mechanisms of breath condensate generation differ significantly depending on physiological differences in lung function, body size, weight and age. Further support of this interpretation comes from the linear correlation of ventilation with total protein in breath condensate. This was true for volunteers as well as for COPD patients although correlation was not as good in the latter group. Alternative, additional or reduced mechanisms of protein entering the cold trap may therefore exist in COPD.

Reinhold et al. have also described a correlation of ventilation (minute volume) and breath condensate volume (per minute) in calves. However, these authors demonstrated a dependency of breath condensate volume on body weight and tidal volume [20]. In our study, lung function parameters were of no significant influence in determining breath condensate volume.

We did not control for a difference in tidal volume in this study in order to avoid any influence on ventilation. Nevertheless, it is suggested from our data that the wide range of volumes ventilated in the 10 min periods (37 – 307 L/min in volunteers and 27 – 238 L/min in COPD) is not merely due to differences in breathing frequency but also involves a range of tidal volumes and nevertheless results in close correlation of...
BCV and $V_E$. The usefulness of breath condensate collection will be greater if tidal volume needs not to be tightly controlled. Our findings do not implicate that tidal volume is a critical factor for neither breath condensate volume nor the constituents protein and urea measured in this study.

The small intra-individual variation in the relation of BCV and $V_E$ and the close correlation of protein with either BCV or $V_E$ suggest that solutes in BCV will also vary only very little within a given healthy subject. In fact, little intra-individual variation between the three collection periods with respect to urea and protein concentration was observed in this series of measurements. Therefore collection of breath condensate is unlikely to be a mechanism that is easily exhausted or altered during the process of breath condensate collection but instead appears to be extendable by increasing the collection period or repetitive collection if more material is needed. We exchanged collecting tubes at 10 min intervals. The capacity of the tubes used in the ECoScreen® appears to be adequate for a thirty minute collection period as was demonstrated by Ct-2 in this study which was removed following a 30 min period instead of 10 and still showed a similar effectiveness in terms of extraction when compared to Ct-1.

In contrast to BCV $V_E$ showed considerable variation as mentioned above. All individuals were asked to breath normally into the mouthpiece of the collection device. Frequently we observed initial short periods of hyperventilation. However five individuals were moderately ($432 \text{ L}, 461 \text{ L}, 464 \text{ L}, 471 \text{ L}, 474 \text{ L}$) and two were markedly ($658 \text{ L}, 850 \text{ L}$) hyperventilating during breath condensate collection. We have intentionally made no effort to influence breathing of volunteers during the collection period in order to test the hypothesis of the linear correlation of ventilation and BCV.

The average volume of breath condensate collected by Ct-1 in our study was 0.95 ml for every 100 L of $V_E$ in healthy volunteers and 1.02 ml for COPD patients. This is slightly below 50% of the entire amount of water in the exhalate calculated from temperature and saturation. When both collecting tubes had been installed in a serial array effectiveness almost doubled. This might be expected since the area for condensation doubled and temperature was very comparable ($-20 \degree C$) in both collecting tubes. However, this demonstrates that in addition to increasing $V_E$, BCV can also be increased by increasing the surface area for condensation even if this means a serial set up. Breath condensate volumes collected from a single trap were similar to the volume collected in 200 cm tubing connected to the expiratory port of a ventilator and running through saline-ice water collecting condensate from mechanically ventilated patients in another study [13].

The relative atmospheric humidity of 88% at the mouthpiece instead of the theoretical humidity of 100% (i.e. BTPS conditions) suggests a dilution of expired gas with air from the nasopharynx. During the passage through the first cold trap humidity fell to an average of 84%. This may not seem to be a large difference, however considering the drop in temperature from 29 to 19 \degree C this relates to a considerable loss of water by condensate formation.

BCV actually collected was approximately 80% of the amount of condensate extracted from the exhalate according to the measurements of humidity and temperature. The residual difference may partially be explained by the loss of small amounts of fluid during recovery from the cold trap (Ct-1, Ct-2) when the frozen condensate has to be thawed and transferred into vials for further processing. The close correspondence of volume extracted and volume actually recovered with some allowance for unrecovered volume also lends support to the interpretation that condensation due to a drop in temperature is in fact the relevant mechanism of breath condensate generation. The alternative possibility of impact driven condensate deposition does not appear to be an important

**Fig. 5** Total protein and urea versus $V_E$. A Linear correlation of total protein with $V_E$ ($r = 0.65$, $p<0.001$, $y = 0.28x - 8.67$). B Linear correlation of total urea with $V_E$ ($r = 0.92$, $p<0.0001$, $y = 0.0054x - 0.44$).
mechanism at least with respect to BCV. If this was untrue, a difference in the ventilation/volume ratio of the first cold trap versus the second cold trap would have been expected, since an impact driven condensate formation would be expected to prefer the first trap.

Protein and urea were both detectable with standard assays in breath condensate. The total amount of protein increased with BCV as well as $V_{E}$ for both groups. For urea this was only true for volunteers, most likely reflecting differences in kidney function of COPD patients. Protein may therefore be a useful denominator for other ELF ingredients e.g. inflammatory mediators.

In conclusion, breath condensate volume is mainly dependent on ventilation, and when corrected for ventilation, is very constant in healthy volunteers as well as in COPD patients. Thus breath condensate volume should be reported with regard to expired volume (rather than time). These observations may help in the interpretation of breath condensate studies and in designing new studies using this indubitably useful tool of pulmonary medicine. As demonstrated here with protein and urea breath condensate contains ingredients most likely derived from the epithelial lining fluid of the lung parenchyma and/or the airways. These ingredients appear to increase linearly with ventilation in healthy individuals and do so albeit more loosely related in COPD patients which suggests that analyses requiring larger amounts of material will be feasible by increasing ventilation or collecting repetitively.

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