Postmortem pulmonary CT in hypothermia

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Abstract So far, fatal hypothermia has been associated with pulmonary edema. With post mortem full body CT scanning (PMCT), lungs can be examined also for CT attenuation. There, low CT attenuation appeared to prevail in lungs of fatal hypothermia cases. Methods. We compared 14 cases of fatal hypothermia with an age-sex matched control group. Additionally, 4 cases of carbon monoxide (CO) poisoning were examined. Furthermore, 10 test cases were examined to test predictability based on PMCT. Two readers measured CT attenuation on four different axial slices across the lungs (blinded to case group and other reader’s results). Results. Hypothermia was associated with statistically significantly lower lung PMCT attenuation and lower lung weights than controls, and there was a dose-effect relationship at a cut-off of 2 degrees C for the environmental temperature. CO poisoning yielded low pulmonary attenuation but higher lung weights. General model based prediction yielded a 94% probability for fatal hypothermia deaths and a 21% probability for non hypothermia deaths in the test group. Discussion. Increased breathing rate is known to accompany both CO poisoning and hypothermia, so this in part could explain the low PMCT lung attenuation due to a ODC (oxygen dissociation curve) left shift. Far more marked distension in fatal hypothermia compared to CO poisoning points to further and possibly different mechanisms. There, increased dead space and increased stiffness to deflation (but not inflation) appear to be effects of cold air (but not CO) that may explain the difference in low PMCT attenuation.
Fig. 1 A low PMCT attenuation in the lungs of hypothermic deaths (bottom row) compared to natural deaths (top row) appeared to be present in fatal hypothermia (but not only there). – Top row: Natural death without signs of hypothermia. Bottom row: Death as consequence of hypothermia. Left column: PMCT image shown using attenuation values that are windowed for bone viewing. Right column: The PMCT images shown that are windowed for lung viewing. All images show axial PMCT images as they are routinely captured before autopsy.

Keywords Computed tomography · hypothermia · postmortem · Virtopsy · Pulmonary emphysema

Key points

– Fatal hypothermia is associated with low PMCT attenuation of the lungs.
– The association between hypothermia and low PMCT attenuation of the lungs exhibits a dose-effect relationship.
– Low PMCT attenuation of the lungs can be found in other situations such as strangulation so it is not specific for fatal hypothermia.
– Macroscopic signs of chronic emphysema were equally frequent in this study’s hypothermia and control group.

Introduction

Diagnosis of fatal hypothermia is based on a number of hypothermia-associated observations as well as on exclusion [1, 2]. At external inspection, bright red
discoloration of livor mortis, red to purple or pink to brownish-pink discoloration over the skin of extensor surfaces such as around elbows or knees, as well as circumstantial indicators such as low environmental temperature or paradoxical undressing may indicate fatal hypothermia. At autopsy, bright red blood, Wischnewski erosions of the gastric mucosa [3], hemorrhages in the psoas muscle and acute pancreatic necroses are among typical [4,5] but not specific [6] findings. Chemical test results that support the assumption of prolonged agony as in fatal hypothermia include ketonuria [7].

PMCT (post mortem full body computed tomography) is currently being included in a number of recent post mortem examination guidelines and standards [8,9]. We routinely perform PMCT scans of bodies admitted to our institute to supplement autopsy. When viewing consecutive full body PMCT scans, it did catch our attention that fatal hypothermia along with a number of other conditions seemed to be associated with a considerably lower PMCT attenuation of the lungs, as opposed to natural causes of death.

Low PMCT lung attenuation typically signifies emphysema. Acute emphysema may be seen in cases of suffocation [10]. With increased breathing rate or obstruction of breathing, conditions such as carbonmonoxide (CO) poisoning or strangulation are expected to correlate with acute agonal pulmonary distension. There was a strong contrast between fatal hypothermia and controls on routine PMCT (see Fig. 1). This was not expected, as forensic pathology texts list pulmonary edema [4,5] or even bronchopneumonia [11] as typical lung findings in such deaths. Conversely, classical forensic pathology books give no indication of there being any correlate for uniformly low PMCT lung

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**Fig. 2 A:** Lungs were analyzed by selecting four axial images - two for each lung with one above and one below the hilus. **B:** Segmentation outlines to define the lung area for CT attenuation averaging were placed manually about 1 cm within the pleura. **C:** We omitted areas of apparent post mortem hypostasis (labeled “H” in the image).
Fig. 3 This box plot shows how CT attenuation (y-axis: Hounsfield Units) differ among the three groups examined: the lung CT attenuation in hypothermic death (middle) is statistically significantly lower compared to controls (right). The CO group (left) also exhibited significantly lower CT attenuation than the controls.

Fig. 4 This image illustrates, on basis of a number of arbitrarily selected axial images displayed at the exact same attenuation-to-grayscale mapping or window, that the lung CT attenuation in hypothermic death (image labels H1 to H4) is lower compared to controls (image labels C1 to C4).

attenuation such as acute emphysema, acute distension or increased agonal hyperventilation in fatal hypothermia.

The aim of this study was to examine PMCT and autopsy correlates of acute distension, acute emphysema and pulmonary edema in cases of fatal hy-
Fig. 5 Logistic regression derived from a General Linear Model shows that given our data (and its inclusion and exclusion criteria), a 90% probability for fatal hypothermia exists for PMCT lung attenuation of less than $-780$ HU, whereas based on this data, the probability for missing fatal hypothermia is less than 10% for PMCT lung attenuation exceeding $-590$ HU. Test data (triangle pointing down: hypothermia, triangle pointing up: not hypothermia) was fitted to test the model, details see Table 2.

...hemia in comparison with controls and in comparison with carbon monoxide poisoning, and to discuss possible explanations for the results.

Materials and methods

We performed a retrospective case-matched study of fatal hypothermia cases (group H) and an age-sex matched control (group C). Additionally, data was also compared with cases of carbon monoxide poisoning (group CO), but since there were just four cases, no age-sex match was possible. Pulmonary PMCT attenuation along with other findings were compared. Last but not the least, a statistical model was determined. The model was then tested with the pulmonary PMCT attenuation of ten additional deaths (group T).

Case review

In our institute, full body PMCT imaging became routine for a number of indications in 2008, and for all bodies that were admitted to our institute in 2010. All cases that had undergone both PMCT and subsequent autopsy at our institute were reviewed for subsequent data collection.
Data collection

Data collection contained the retrospective identification of data and protocols that had been, in their entirety, performed under the auspices of investigatory authorities for judicial investigations into suspicious or violent deaths. Selection criteria for the three groups (H, C, CO) are detailed below. Data was collected in accordance with our institute’s policies, local ethical committee’s opinion and laws pertaining to biomedical research.

Data analysis

Autopsy data and circumstantial evidence were evaluated by one author of this study. From the autopsy reports, age, sex, temperature at death scene, post mortem interval, and other details including the presence of pulmonary edema and lung weights were obtained. PMCT data was also evaluated for presence of gas. PMCT attenuation was measured by two other authors as blinded readers.
Fig. 7 This plot shows modeled curves for \(O_2\)-Hb dissociation curves (ODC) under influence of both carbon monoxide poisoning and cold temperature.

Hypothermia group (H)

We were interested in pulmonary findings that resulted from direct influence of hypothermia before and during agony and death. Therefore, complications of initially survived hypothermia were not evaluated in this study.

A 14 deaths containing all cases where (dry) hypothermia [4] was judged to be the main (or a contributing factor to the) cause of death resulted from the case review. There were no cases where findings of possible hypothermia were identified alongside drowning. There were 5 women with a mean age of 55 ± 21 years and 9 men with a mean age of 54 ± 22 years.

Further diagnostic considerations regarding fatal hypothermia are detailed below.

Control group (C)

The control group was age-sex matched to group H. Cases were selected randomly from the case review’s respective age-sex-group (that was defined as age matching the age of the respective individual of group H ±2.5 years and same sex).

Excluded mechanisms contained drowning, gaseous and mechanical asphyxia such as strangulation and carbon monoxide poisoning. Advanced post
mortem decomposition cases were excluded. In these cases, morphological assessment of possible fatal hypothermia was too uncertain. Also, decomposition related gas was assumed to possibly confound possible effects of hypothermia on the PMCT attenuation of lungs.

Carbonmonoxide poisoning group (CO)

From the case review, all cases of fatal carbonmonoxide (CO) poisoning were included for this study in order to investigate the effect of suffocation on PMCT appearance of the lungs. The defining criterium for selecting these was whether carbonmonoxide was judged to be the cause of death in the final report of the pathologist; carbonmonoxide-hemoglobin values were relatively high (see Table 1).

We omitted cases of advanced post mortem decomposition and cases with competitive causes of death such as heat or fire consummation, traumatic injury or hyperthermia. No age-sex match to the other groups was performed due to the small group size.

Diagnostic considerations

Circumstantial evidence that was interpreted as diagnostic of hypothermia contained a relatively low environmental temperature. In this study, environmental temperature in group $H$ was $9.8 \pm 8.6 \, ^\circ C$ on average, compared to the control group's mean of $22.0 \pm 2.2 \, ^\circ C$ (statistically significant, Wilcoxon $p<0.001$).

Autopsy findings to suggest fatal hypothermia contained livid skin discoloration over elbows and knees (8/14 in group $H$), gastric Wischnewski type erosions (9/14 in group $H$), hemorrhages within the Psoas muscles (2/14 in group $H$) and a positive qualitative test for ketone bodies in the urine (9/14 in group $H$). Pancreatic necroses had not been found in these cases. In no deceased of group $H$ had there been any indication of positional or other mechanical asphyxia.

Gas due to putrefaction may also contribute to a low lung PMCT interpretation. So we documented the presence of gas as found in post mortem decay [12] or after trauma [13]. PMCT data was evaluated for gas in the heart, pulmonary vessels, liver vessels and subcutaneous soft tissue on the post mortem CT scans (one examiner, semiquantitative evaluation, see Table 1). Post-mortem interval was around 32 – 36 hours on an average for each of the three groups $H$, $C$ and $CO$.

See table 1 for an overview.
Ten additional cases were selected, using the selection criteria outlined above for group H (n=7) or C (n=3). Their pulmonary PMCT attenuation results were exclusively used to test (not to determine) the statistical model.

The seven cases of fatal hypothermia contained gastric erosions (5/7), livid skin discolorations (5/7), a positive qualitative test for ketone bodies in the urine (5/7) and psoas hemorrhages (2/7).

For more details of group T, see Table 2.

Lung PMCT attenuation value measurement

Data collection of PMCT attenuation of the lungs was performed as a two reader study. Both readers had considerable experience in PMCT image reading, reporting and validation of findings at autopsy.

Groups H, C, CO and T contained no deceased with less than two complete lungs. For the left and right lung and for both the suprahilar and infrahilar level as depicted in Figure 2, axial images were selected by the first examiner (R1, radiologist). In total, four axial lung PMCT images were used to evaluate each case. Serial numbers of axial images that were selected by the first reader were documented. With these serial numbers, the second reader (R2, forensic pathologist) could analyze the same.

The first reader defined the slices of the axial images and performed attenuation measurements. The second reader used, for each case, the same slice axial images previously examined. The second reader performed evaluations on another computer and at another time and independently. Lung cross sectional areas were first segmented manually on standard axial images (Figure 2): a marker line was placed about 1 cm within the perimeter of the pleura. Then, an average CT attenuation value was obtained using standard software; results are given in Hounsfield units (HU). Areas of what appeared to be marked pulmonary hypostasis (internal livor mortis of the lung [14], a typical post mortem finding) were not included in our segmentation. The reason is explained below. Determination of PMCT values was performed in a no particular order. Examiners were blinded to case details and the other examiner’s findings. For each case, results were averaged before they were compared between observers in order to yield observer agreement. For each case and across both observers, results were averaged to yield one CT attenuation value.

Presence of pulmonary hypostasis

Pulmonary hypostasis is regarded as a phenomenon of gravity dependent accumulation of blood [14]. As a predominantly radiological organ finding similar to livor mortis of the skin, it is usually not examined or described at autopsy. Pulmonary hypostasis is not thought to be related to physiological ante mortem
or agonal processes. Therefore, we decided to exclude pulmonary hypostasis from our PMCT attenuation measurements as much as practically possible.

Nevertheless, presence and degree of hypostasis were documented. Presence of hypostasis was judged subjectively. Its visible presence was noted for each case (see Table 3). In addition, difference between CT attenuation was determined by selecting highest and lowest lung attenuation. The highest lung attenuation values were mostly dorsally located, the lowest anteriorly. Measurements were performed by capturing the average attenuation of an 8 – 10 cm² region using a region-of-interest measurement tool (see Table 3).

**Statistical considerations**

**Reader agreement:** Reader agreement between both readers was determined using the Intra-Class Correlation coefficient (ICC [15,16]).

**Comparison of PMCT attenuation:** A Shapiro-Wilk test [17] for each of the age-sex matched groups’ PMCT attenuations (group C, group H) yielded a statistical significance of p< 0.01. So, the normal distribution assumption was rejected. Results for the CO group conformed to this test. Variances of all three groups H, C and CO were different according to Bartlett’s K-squared test of homogeneity of variances [18]. With that, we chose Wilcoxon’s nonparametric test [19] over Student’s t for statistical comparison of PMCT attenuation values (Figure 3).

Lung PMCT reading, qualitative appreciation and check of plausibility was performed by applying the same attenuation-to-grayscale windowing settings to axial images of all groups (Fig. 4).

**Logistic regression:** Our data yields a sparseness indicator of $\psi < 0.1$ for the attenuation data of both groups H and C (see Equation 1 quantifying sparseness for vector $v[1,..,n]$ with $\psi = 0$ for dense and $\psi = 1$ for maximally sparse vectors which uses an $\ell_1/\ell_2$-norm penalization [20]). This indicates however that Wald statistics are applicable in this instance [21] as no Hauck-Donner effect should be expected. Subsequently, a General Linear Model (GLM) ([22]) assuming a binomial data distribution to determine membership to either H or C (equal group sizes) was employed in order to perform and plot [23] logistic regression (Figure 5).

Determination of probability for group T members to be either hypothermic or not hypothermic was achieved by inserting lung PMCT attenuation data of T into the fitted GLM.

$$\psi(v) = \sqrt{n - \| v \|^2} = \frac{\sqrt{n} - \frac{\sum_{i=1}^{n} v_i}{\sqrt{\sum_{i=1}^{n} v_i^2}}}{\sqrt{n - 1}} (1)$$

**Dose-effect correlations – temperature dependent lung attenuation differences within group H:**
We were interested to find out whether there may be a cause-effect relationship. With that assumption and within group H, lower environmental temperatures would have to correlate with even lower PMCT lung attenuation. To investigate such a possible dose-response relationship between environmental temperature and PMCT lung attenuation, the hypothermia group H was split into two groups based on temperatures at death scenes. This was done along a range of different cut-off values $T_C \in [1,7]^\circ C$ for the environmental temperature $T$.

We obtained a number of dichotomous subgroups of $H$ of different sizes and variances as a function of the continuously varied value of $T_C$: one hypothermia group is characterized by a lower environmental temperature $H_L$, the other group $H_H$ characterized by higher environmental temperature, with $H = H_L \cup H_H$ and environmental temperatures $T$ conform to $T_{H_L} < T_C \leq T_{H_H}$.

We then compared statistical properties of PMCT attenuation within $H_L$ and $H_H$. With variance unequality and non-normal data distribution being erratically distributed across the continuity of the varied cut-off values $T_C$ (Shapiro Wilk’s test for normality and Bartlett test for equal variances, see Fig. 6), we used the non-parametric Wilcoxon test to examine statistical significance in PMCT lung attenuation.

And so with varying $T_C$, at a cut-off value of $T_C = 2^\circ C$, a split of group H into $H_L$ and $H_H$ yielded statistical significance for PMCT lung attenuation (Figure 6).

**CO-Hb dependent lung attenuation differences within group CO:** In the same manner as splitting group H based on environmental temperature, group CO was split based on CO-Hb concentrations. A best cut-off of 70% CO-Hb resulted, however due to low group member count, only numerical results (but no statistical evaluation) will be reported.

**Contingency tables:** For statistical comparison against the other groups, Barnard’s exact test was used (CSM [24–26]).

Mathematical model of the effect of carbon monoxide and hypothermia on the oxygen dissociation curve (ODC)

A carbon monoxide-hemoglobin (CO-Hb) concentration dependent left shift of the oxygen dissociation curve (ODC) results in a pathologically increased hemoglobin’s $O_2$-saturation. Increasing CO-Hb concentrations yield an increasing disturbance of $O_2$ delivery from capillaries to the tissues [27]. CO poisoning as well as the effects of acute hypothermia [28] both cause a similar left shift of the ODC. To compare their magnitude, we implemented a combined algorithm in R [22]: one approach models the influence of elevated CO-Hb concentration on the ODC shift, and the other one models the increasingly low temperature on the ODC left shift[29,30]. We compared our resulting curves with published diagrams from authors covering both temperature [31] and CO-Hb [32] effects. That comparison appeared to plausibly validate the combined algorithm.
Whole body CT scans were obtained on a standard scanning device (Somatom Definition Flash, Siemens, Erlangen, Germany). Scans were obtained at 120 kV, reference mAs 400 and 128 x 0.6 mm collimation with automated dose modulation (CARE dose4D, Siemens, Erlangen, Germany) [33]. Slice thickness was 2 mm with an increment of 1 mm. Axial slices were reconstructed with a relatively hard kernel (“B60f”, Siemens, Erlangen, Germany). Manual lung cross section segmentation and CT attenuation measurements were performed using standard radiology image viewing software (Sectra Workstation IDS7, Version 14.3.9.3, Sectra Imtec AB, Linköping, Sweden). Statistical evaluation was performed using R (R-package[22] Version 2.12.1, including add-ons for intra class correlation coefficient (IRR [16] Version 0.83) and Barnard’s exact test (Exact package[25] Version 1.4)) and JMP (Version 9, SAS Institute, Cary NC, USA).

Results

PMCT lung attenuation reader agreement

This was done as a two reader study. Reader agreement of pulmonary PMCT attenuation determination was good with an intra-class correlation coefficient of 0.86.

Pulmonary hypostasis

Pulmonary hypostasis was significantly less frequently present and significantly less pronounced in fatal hypothermia cases compared to controls or CO poisoning (see Table 3).

PMCT lung attenuation, lung weights and pulmonary edema

PMCT lung attenuation was assumed to correlate with both autopsy findings of pulmonary edema and increased lung weights. This bases on the assumption that attenuation of air is higher than attenuation of water. In order to examine this assumption, PMCT lung attenuation was correlated with the diagnosis of pulmonary edema made at autopsy and with lung weights.

– In cases with the autopsy diagnosis of pulmonary edema (across all three groups \( H, C, CO \)), PMCT lung attenuation was significantly \( (p < 0.03, \) Wilcoxon) higher \( (−619 ± 92 \text{HU}) \) than in cases with no autopsy finding of pulmonary edema \( (−726 ± 167 \text{HU}) \).
Group size | Group H | Group C | Group CO
--- | --- | --- | ---
14 | 14 | 4
Age | 54.5 ± 20.5 y | 54.4 ± 20.5 y | 55.8 ± 8.7 y
Sex | 5 female / 9 male | 5 female / 9 male | 2 female / 2 male

| Age | 54 | 54 | 35 |
| ± | ± 20 | ± 20 | ± 8 |
| y | y | y |

| Sex | female | male | female | male |
| | | | | |

| Temperature at death scene [°C] | 9.8 ± 8.6² | 22.0 ± 2.2² | 18.1 ± 19.4 |
| Post mortem interval to CT [h] | 36 ± 33 | 32 ± 24 | 34 ± 31 |
| Acute intoxication (alcohol and / or drugs) | 7/14 | 1/14 | 1/4 |
| CO-Hb [%] | n.r. | n.r. | 73 ± 10 |
| Trauma | 5/14 | 2/14 | 1/4 |
| Spontaneous intracranial hemorrhage | 2/14 | 1/14 | 0/4 |
| Coronary artery atherosclerosis | 2/14 | 8/14 | 0/4 |
| Sudden death, epilepsy | 0/14 | 1/14 | 0/4 |
| Acute myocardial infarction | 1/14 | 5/14 | 0/4 |
| Chronic alcoholism | 4/14 | 0/14 | 0/4 |
| Lung weight [g] | 1242 ± 504² | 1599 ± 524³ | 1557 ± 433 |
| Pulmonary edema | 3/14⁴ | 10/14⁴ | 3/4 |
| Gastric erosions | 9/14⁴ | 3/14⁴ | 0/4 |
| Chronic pulmonary emphysema | 3/14⁴ | 4/14⁵ | 0/4⁶ |
| Gas (pulmonary vessels) | no 14/14 | no 14/14 | no 4/4 |
| | present 0/14 | present 0/14 | present 0/4 |
| Gas (soft tissue) | no 14/14 | no 13/14 | no 4/4 |
| | present 0/14 | present 1/14 | present 0/4 |
| Gas (liver) | no 13/14 | no 5/14 | no 4/4 |
| | minimal 1/14 | minimal 7/14 | minimal 0/4 |
| | extensive 0/14 | extensive 2/14 | extensive 0/4 |
| Gas (heart) | no 13/14 | no 7/14 | no 4/4 |
| | minimal 1/14 | minimal 5/14 | minimal 0/4 |
| | extensive 0/14 | extensive 2/14 | extensive 0/4 |

Table 1

Group H (hypothermia) differs from the control group (C) in a number of details. Group H, both due to absence of resuscitation measures and extensive trauma, also contains considerably less gas.¹ Age-sex matched groups, differences statistically significant with ²p < 0.001 (Wilcoxon), with ³p < 0.03 (Wilcoxon), with ⁴p < 0.004 (Barnard’s exact test) and with ⁵p < 0.03 (Barnard’s exact test) and ⁶no significant difference (H vs. C vs. CO, Chi-Square).

– Lung weights were correlated with PMCT lung attenuation across the whole collective (all groups H, C and CO). Least squares regression established a positive correlation, analysis of variance yielded a significance level of p < 0.001 (F-ratio of 14.2).

As hypothermia appears to entail markedly distended or fluid deprived lungs based on PMCT appearance, we compared lung weights and presence of pulmonary edema. Autopsy derived lung weights (H: 1242 ± 504g, C: 1599 ± 524g, Wilcoxon p < 0.03; CO: ) and pulmonary edema diagnoses (H: 3/14 cases, C: 10/14 cases, Barnard’s exact test p < 0.004) differed significantly be-
Fatal hypothermia  |  No hypothermia  
---|---
Group size  | 7  | 3  
Temperature at death scene [°C]  | 8.5 ± 6.1  | 20.0 ± 2.0  
Acute intoxication (alcohol, drugs)  | 2/7  | 0/3  
Coronary artery atherosclerosis  | 2/7  | 1/3  
Chronic pulmonary emphysema  | 1/7  | 0/3  
Trauma  | 1/7  | 0/3  
Acute myocardial infarction  | 0/7  | 1/3  
Brain stem hemorrhage  | 0/7  | 1/3  
Pneumonia  | 0/7  | 1/3  
PMCT lung attenuation [HU]  | −822 ± 43  | −564 ± 121  
GLM predicted probability of membership to group H  | .94 ± .04  | .21 ± .34  

### Table 2 Test group details.

<table>
<thead>
<tr>
<th>Hypothermia</th>
<th>CO poisoning</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulmonary hypostasis presence</td>
<td>2/14 (14%)</td>
<td>4/4 (100%)</td>
</tr>
<tr>
<td>Pulmonary hypostasis - difference in CT attenuation [HU]</td>
<td>91 ± 76</td>
<td>496 ± 78</td>
</tr>
<tr>
<td>PMCT lung attenuation [HU]</td>
<td>−762 ± 66</td>
<td>−695 ± 65</td>
</tr>
<tr>
<td>Autopsy lung weights [g]</td>
<td>1242 ± 504</td>
<td>1557 ± 433</td>
</tr>
<tr>
<td>Pulmonary edema</td>
<td>3/14</td>
<td>3/4</td>
</tr>
<tr>
<td>Increased breathing rate</td>
<td>++</td>
<td>++</td>
</tr>
<tr>
<td>Dose dependent ODC left shift</td>
<td>++ (Fig. 7)</td>
<td>++ (Fig. 7)</td>
</tr>
<tr>
<td>Dead space</td>
<td>+[34]</td>
<td>+[35]</td>
</tr>
<tr>
<td>Cold temperature induced lung stiffness</td>
<td>+[36,37]</td>
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</tr>
</tbody>
</table>

### Table 3 Summary of results and interpretation.

Hypostasis was not diagnosed in most cases of fatal hypothermia with a significantly lower position dependent lung attenuation difference compared to both CO and C groups (significance ¹ p<0.0001 Chi-Square, ² p<0.0001 Wilcoxon). While both fatal hypothermia and carbon monoxide poisoning exhibit significantly reduced PMCT lung attenuation, lower lung weights in fatal hypothermia compared to CO poisoning indicate that mechanisms that differ between both noxious agents need to be considered.

Between the hypothermia (H) and control (C) groups. So, hypothermia was less frequently associated with pulmonary edema than the control group, and lung weights in fatal hypothermia were significantly lower. Average lung weights in carbon monoxide poisoning (CO: 1557 ± 433g) were higher than the hypothermia group’s lung weights (Wilcoxon with no statistical significance); pulmonary edema was diagnosed significantly more in carbon monoxide poisoning (CO: 3/4 cases) than in the hypothermia group (H: 3/14 cases; p< 0.04, Barnard’s exact test).
Averaged lung PMCT attenuation was $-762 \pm 66$ HU in group H and with that, it was significantly lower than the mean attenuation of $-546 \pm 152$ HU in group C ($\text{Wilcoxon } p < 0.001$). PMCT lung attenuation of the CO group was $-695 \pm 65$ HU and with that, it also exhibited significantly decreased attenuation compared to controls ($\text{Wilcoxon } p < 0.03$), but higher than group H, (Wilcoxon test: difference between groups CO and C not statistically significant) (see Figure 3).

These PMCT attenuation differences between fatal hypothermia and controls are also visibly plausible. This becomes apparent when comparing hypothermic (H1 through H4, Figure 4) and control (C1 through C4, Figure 4) lungs in PMCT using the same reconstruction kernel and the same windowing.

Predicting fatal hypothermia

Fatal hypothermia may be suspected based on above listed criteria and PMCT findings in a new case. To test that assumption, ten test cases (group T) were evaluated using the GLM (i.e., the statistical model) that had been determined based on existing data of groups H and C.

The logistic regression curve (see Figure 5) derived from the GLM shows a significant statistical effect of pulmonary PMCT attenuation on group membership H or C (given their group definitions), respectively (Wald statistic: $p< 0.001$). The GLM also predicted that given our test data (and its inclusion and exclusion criteria), there would be a 90% probability for hypothermia with PMCT lung attenuation of less than $-780$ HU, whereas the probability for overlooking hypothermia is less than 10% for PMCT lung attenuation greater than $-590$ HU.

The seven hypothermic cases of test group T were predicted to be hypothermic based on PMCT lung density alone, with a probability of 94% (see Table ??). The three control cases of group T yielded an average fatal hypothermia probability of 21%.

To illustrate the test results given the model, test case CT densities were plotted alongside the fitted model data as triangles (see Fig. 5).

Dose-effect correlations: temperature dependent lung attenuation differences within group H

We investigated whether our data contained correlations indicative of a dose-response effect. Below 2 °C, PMCT yielded lung attenuation of $-840 \pm 44$ HU, whereas environmental temperatures $\geq 2$ °C correlated with lung attenuation of $-749 \pm 62$ HU ($\text{Wilcoxon } p < 0.02$).

So lower temperatures correlated with lower PMCT lung attenuation than higher temperatures within the victims of fatal hypothermia.
Dose-effect correlations: CO-Hb dependent lung attenuation differences within group CO

The group of fatal CO-poisoning contained 4 deaths. At a cut-off of 70% CO-Hb, lung attenuation averaged $-672 \pm 108$ HU in the lower CO-Hb group (mean 58% CO-Hb) compared to a slightly lower pulmonary attenuation in the higher CO-Hb group (mean 78% CO-Hb) of $-702 \pm 47$ HU.

So higher CO-Hb concentrations correlated with lower lung attenuation in PMCT.

Comparison between the effects of hypothermia and carbon monoxide poisoning on the oxygen-hemoglobin dissociation curves (ODC)

The direct effect of carbon monoxide (CO) poisoning and hypothermia on the ODC were compared on a one to one basis using a curve plot we obtained from implementing a combined algorithm. With that, a 30% CoHb concentration has a comparable effect on the ODC as a temperature of 29°C, and comparable values to increasing values of CO poisoning and hypothermia pair 60% CoHb/22°C and 75% CoHb/9°C body core temperature (see Figure 7).

Discussion

We found pulmonary attenuation in PMCT to be significantly lower in fatal hypothermia than in age-sex matched controls. This was not due to post mortem gas accumulation as consequence of trauma, resuscitation or putrefaction. Much rather, the study results suggest that this is an effect of fatal hypothermia.

Lung attenuation of less than $-780$ HU appeared to correlate with an observed probability of hypothermic death exceeding 90% (excluding drowning, gaseous and mechanical asphyxia such as strangulation and carbon monoxide poisoning). A predictive probability derived from Generalized Linear Model that we determined from study groups H and C placed a 94% probability on seven new test cases of fatal hypothermia compared to a 21% probability for three non hypothermic deaths in the test group. A number of studies that were published [38,39] in parallel while first results of our study had appeared in conference proceedings ([40], online since August 1st 2012) seem to confirm our findings of decreased PMCT pulmonary attenuation setting off fatal hypothermia against other cases.

The question is what that observation means.

Lower than normal attenuation of lungs in CT usually is caused by overinflation or distension. In the living, emphysema is associated with CT attenuation of less than $-950$ HU [41,42]. Normal clinical lung attenuation ranges from $-797$ HU (10% vital capacity, expiration) to $-877$ HU (90% vital capacity, inspiration) [43]. Conversely, significant pulmonary edema is primarily regarded
as accumulation of fluid in airspaces [44]. Presence of pulmonary edema will increase PMCT attenuation of the lungs as shown on PMCT scans; typical CT-appearance of significant edema also is described as ground-glass opacity [45].

In our data, increased PMCT attenuation of lungs correlates with autopsy findings (increased lung weight, pulmonary edema). Conversely, low PMCT attenuation correlates with absence of pulmonary edema and normal autopsy lung weights. Average weight of normal autopsy lungs is reported as 1246 g [46]. So the average lung weight of 1242 ± 504g in our hypothermia collective did not suggest an increase of fluid content over normal lungs.

Pulmonary edema generally is prevalent in natural death, particularly in conjunction with heart disease [47]. Conversely, pulmonary edema is also seen after intoxication, such as heroin [48–50] or alcohol intoxication. In our control group, lungs weighed 1599 ± 524g on an average and PMCT lung attenuation was −546 ± 152 HU. Textbooks covering lung findings in fatal hypothermia mention pulmonary edema [4,5,11], but that was not a finding that we found to be typical. However, low PMCT attenuation (average of −672 HU) also was observed in four cases of CO poisoning. There, autopsy had yielded increased lung weights (average of 1557 ± 433g) and pulmonary edema.

So, in our data, fatal hypothermia is associated with distended lungs without signs of pulmonary edema. Carbon monoxide poisoning correlates with distended lungs or acute emphysema, alongside pulmonary edema.

Findings of acute pulmonary emphysema in forensic pathology can be typically found in strangulation [51]. Dry and lightweight lungs are also known from ‘typical’ drowning cases particularly in fresh water [52]. In drowning, dry lungs may be seen only within a relatively short post mortem interval, whereas bodies retrieved from water after two days or more generally tend to exhibit heavier lungs [53]). Attempts for forced breathing or increased breathing may occur in strangulation as well as in drowning. To explain increased breathing rates in hypothermia or in carbon monoxide poisoning [54], the left shift of the ODC could be a plausible model. Severe hypothermia and carbon monoxide both have a direct effect on the ODC, and the magnitude of these effects can be compared (see Figure 7).

However, the degree of pulmonary dryness and distention in fatal hypothermia exceed that in CO poisoning, so further mechanisms may have to be considered.

Cold temperatures also cause an increased resistance to deflation: a 34% decrease in dynamical compliance was observed to be due to cold-dependent mechanical changes of the lungs [36]. At relatively low lung tissue temperatures such as 4 °C, deflation was found to require more pressure than inflation [37]. Muscular energy dies down in the process of hypothermic agony. Together with rapid cooling rates, forces required to inflate the lungs then might still be provided by the weakened muscles, when the same muscles may not be as well able to deflate these lungs as well any more as higher forces are required for that. This combination then may contribute to yield an inflated appearance despite relative reduction of tidal volumes being breathed.
In addition, cold temperatures can increase dead space, entailing a striking absence of fluid normally contained within pulmonary blood vessels. Hypothermia is known to massively reduce pulmonary perfusion, and even at around 29 °C, over half of the lung capillaries may already be shut down [34]. Conversely, CO poisoning does not seem to increase dead space by that much [35].

We found a dose-effect relationship in hypothermic deaths (study group H), whereas bodies found at environmental temperatures below 2°C exhibit a statistically significantly lower PMCT lung attenuation than bodies found at or above 2°C. Furthermore, slower body cooling rates were found to differently impact other aspects of metabolism [55] compared to faster cooling rates.

A dose-effect relationship could also be assumed from a difference within the CO group. Physiological tests and mathematical models document an increase in breathing volume with higher carbon monoxide levels [56].

Given these results and interpretation, low PMCT lung attenuation may indicate fatal hypothermia under the entry criteria for both groups H and C. Even for the purpose of conservative forensic pathology where X-ray based examinations are restricted [57] or entirely avoided [58], diagnostic criteria that identify fatal hypothermia while examining a cause of death may be of interest. This even more so in the context of a mountain environment, where hypothermia can be hard to diagnose when people may die at low temperatures at relatively high altitudes, and the body then is retrieved for post mortem examination in the midlands.

Limitations of this study

Low PMCT attenuation of lungs might be a relatively sensitive finding for hypothermia. Also, CO poisoning was found to correlate with below-normal PMCT lung attenuation in our data. So we have to assume that it is not very specific. This study is further limited in that cases of putrefaction, suffocation including drowning (and in particular, drowning in cold water) and strangulation were excluded.

Case count in this study is relatively low. Given our specific environment and its climate, fatal hypothermia typically contains relatively low temperatures compared to other countries [59]. Our hypothermia group thus might be limited to cases with rather rapid agony, whereas pulmonary edema or pancreatic necroses might be findings pertaining to more extended agony intervals. Some well-known vital reactions to hypothermia may take more time to develop, such as pancreatic necroses, and these were not found at autopsy in any of our cases.

Conclusions

Pulmonary PMCT attenuation in fatal hypothermia as well as autopsy lung weights was found to be significantly lower compared to age-sex matched con-
trols. However, we excluded cases with advanced decomposition from the hypothermia group and we excluded a number of diagnostic groups from the control group.

Underlying mechanisms behind this difference might be found in a significant left shift of the ODC. Other factors that were identified as specific for hypothermia include a temperature dependent increase of deflation over inflation pressures in the hypothermic lung as well as marked increase in dead space. The PMCT finding of strikingly low lung attenuation can be helpful for screening purposes. There, causes of death such as suffocation including CO poisoning, suffocation or strangulation have to be considered, but also, fatal hypothermia should be investigated.

**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**


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