A fatal case of oxygen embolism in a hospital

Lionel Commenta, Vincent Varletb, Kewin Ducrotc and Silke Grabherrc

aForensic Medicine Unit, University Center of Legal Medicine, CHUV Hospital, Lausanne-Geneva, Switzerland; bForensic Toxicology and Chemistry Unit, University Center of Legal Medicine, CHUV Hospital, Lausanne-Geneva, Switzerland; cUnit of Forensic Imaging and Anthropology, University Center of Legal Medicine, CHUV Hospital, Lausanne-Geneva, Switzerland

ABSTRACT
This case reports on a 68-year-old man who was found dead in hospital next to his bed. Before this, he had been treated with intravenous antibiotics for pneumonia. The body was found with a peripheral venous catheter connected to a nasal cannula delivering oxygen (O₂) from the wall. Extensive medicolegal examinations were performed, including post-mortem computed tomography (CT), complete conventional autopsy, histological and immunohistochemistry analysis, toxicological analysis and post-mortem chemistry. Additionally, CT-guided gas sampling was performed at multiple sites to collect samples for gas analysis. During the external examination, massive subcutaneous emphysema was visible over the entire surface of the body. The CT scan revealed the presence of gas throughout the vascular system, and in the subcutaneous and muscular tissues. The autopsy confirmed the presence of lobar pneumonia and multiple gas bubbles in the vascular system. The gas analysis results showed a subnormal concentration of oxygen, confirming the suspected pure O₂ embolism. Moreover, the carbon dioxide (CO₂) concentration in the gas sample from the heart was elevated to a level similar to those found in scuba diving fatalities. This could come from degassing of dissolved CO₂ that accumulated and was trapped in the cardiac cavity. Based on the results of the different exams performed, and especially the gas analysis results, it was concluded that the cause of death was O₂ embolism.

Introduction
In addition to autopsy, many tools can be used by a forensic pathologist to determine the cause of death. These tools include post-mortem imaging, toxicological analysis, post-mortem chemistry, histology and immunohistochemistry. Although none of these analyses can replace autopsy, they all can help to discriminate between several possible causes of death and confirm the results obtained from other techniques. When available, the circumstances of the death, and the medical and family history of the individual can help the forensic pathologist to perform the autopsy and choose the most useful complementary exams.

Nowadays, post-mortem imaging is very well developed [1,2], and a contrast agent can be used to obtain a very detailed view of the vascular system [3,4]. Gas accumulated in the body can be precisely located and quantified using post-mortem imaging [5], and an assessment of any post-mortem changes [6] can be made by a forensic radiologist to differentiate between an exogenous or an endogenous (i.e. putrefactive) origin of the gas [7]. Analysis of the gas to determine its composition can confirm the imaging findings and help to differentiate between endogenous and exogenous origins [8]. Gas analysis results can indicate if the gas is from air embolism or physiological changes occurring during the perimortem phase [9,10].

At the University Center of Legal Medicine in Lausanne, Switzerland, autopsies are performed at the request of a prosecutor. A native computed tomography (CT) scan is routinely performed before the autopsy, and, in the case of a suspected vascular lesion, a multi-phase post-mortem CT angiography is performed [11]. Samples of major organs (brain, heart, lungs, liver and kidneys) are taken during the autopsy for histological analysis [12]. Multiple samples are taken for toxicological analysis, post-mortem chemistry and genetic analysis. In some specific cases involving gas poisoning, or if unexpected accumulations of gas are identified after the native CT scan, gaseous samples are also taken according to the protocol proposed by Varlet et al. [7].

The present case describes a fatality following a (probable) suicidal oxygen (O₂) embolism at a hospital and demonstrates the usefulness of gas sampling and analysis for precise diagnosis of gas embolism. To the best of our knowledge, this is the first case described in the literature of CT-guided gas sampling and analysis for a fatal O₂ embolism.
Case report

A 68-year-old male was admitted to hospital with a history of dyspnea at rest for 2 d, a cough producing mucus, loss of appetite and diarrhoea. Pneumonia was suspected, and the patient received intravenous antibiotic treatment. During the patient’s second night in hospital, he was found groggy and walking near his room by the nurses. They took him back to his room and connected a new antibiotic dose to his venous catheter. A few hours later, he was found unconscious, lying on the floor next to his bed. His peripheral venous catheter was connected to a nasal cannula delivering O₂ from the wall, with a flow rate of approximately 2 L/min, and the antibiotic dose was found in the garbage can. As post-mortem lividity was present, no resuscitation attempts were performed by the clinicians and the death was pronounced. The body was immediately brought to the University Center of Legal Medicine (Lausanne, Switzerland).

CT imaging and gas sampling

Before any manipulation of the corpse, a native CT scan was carried out at around 10 h post-mortem using an eight-row CT unit (CT LightSpeed 8, GE Healthcare, Milwaukee, WI). All scanning parameters are detailed in Table 1. A forensic pathologist immediately viewed the native CT images. To evaluate the distribution of gas because of physiological changes in the body after death, the radiological alteration index (RAI) was used as proposed by Egger et al. [6]. The RAI was based on the analysis of samples from seven sites (heart cavities, liver parenchyma and vessels, left innominate vein, abdominal aorta, kidney parenchyma, L3 vertebra and the subcutaneous pectoral tissues).

All images were interpreted in a consensus reading by one board-certified radiologist and one forensic pathologist who were trained in forensic imaging. A post-mortem radiological report was prepared and described all findings from the native CT scan. The native CT scan revealed the presence of subcutaneous, intramuscular and intravascular gas, a pneumoperitoneum, a bloated heart with the right cavities filled with gas and a left pneumothorax (Figure 1(a–d)). The maximum RAI score is 100, and RAI scores greater than 50 are usually seen in cases of severe changes [6]. Therefore, given the short post-mortem interval (<12 h), the elevated RAI obtained in this case (75) strongly suggests an exogenous source of gas in the tissues.

CT-guided gas samples were immediately taken from multiple sites (carotid artery, jugular vein, right auricle, thoracic aorta, pectoral muscle, thoracic cavity, abdominal cavity, scrotum and gluteal soft tissues, Figure 1(e,f)) according to the protocol for gas analysis described by Varlet et al. [7].

External examination

The external examination revealed massive subcutaneous emphysema with audible crepitations on the entire surface of the body. Additionally, there was a bruise on the left part of the forehead, a small contusion near the left eyebrow surrounded by a purplish-blue bruise and some bruises of different ages on the inferior part of the thorax and on the legs. On the left arm, a venous catheter was still in place.

Toxicological and biochemical analysis

Samples for toxicological and biochemical investigations (blood and urine) were collected in S-Monovette® tubes with sodium fluoride or ethylenediaminetetraacetic acid as a preservative (Sarstedt, Nümbrecht, Germany). Biological samples were collected as soon as possible on arrival of the body at the morgue (vitreous humor) and during autopsy (femoral blood, pericardial fluid and urine). During sampling of femoral blood by incision with a scalpel, multiple gas bubbles were visible in the blood (Figure 2(a)).

Toxicological analysis of the collected samples revealed acetone at a physiological level in the blood, as well as caffeine and paracetamol in both blood and urine. Post-mortem chemistry analysis of the serum

| Anatomical region                  | Scan type   | Thickness slice table speed pitch | Interval spacing | Scan field of view (FOV) | Kilovolts (kV) | Milli amperage (mA) | Algorithm of reconstruction |
|-----------------------------------|-------------|----------------------------------|------------------|--------------------------|---------------|---------------------|---------------------------|
| Brain                             | Axial 2.0 s | 2.5 mm (base) 5.0 mm (top)       | 2.5              | 25                       | 120           | 300                 | Standard                  |
| Skull/brain/neck                  | Helical 1.0 s | 1.25 mm 13.50 1.35:1            | 1                | 25                       | 120           | 100–300              | Standard/bone             |
| Thorax/abdomen                    | Helical 1.0 s | 1.25 mm 13.50 1.35:1            | 1                | 50                       | 120           | 150–300              | Standard/bone             |
| Lower extremities                 | Helical 0.8 s | 1.25 mm 13.50 1.35:1            | 1                | 50                       | 120           | 100–220              | Standard/bone             |
| Gas sampling protocol (scan for setting needle punctures) | Helical 0.8 s | 5 mm 33.5 1.675:1               | 5                | 50                       | 120           | 80–200               | Standard                  |
Figure 1. Images obtained by native post-mortem CT showing massive accumulation of gas in all anatomical compartments. (a) Presence of gas in the thorax, including gas in the cardiac cavities (CC), the pleural cavities (PC), the abdominal cavity (AC) and the aorta (A). (b–d) Axial images obtained at the level of the head (b), the neck (c) and the pelvis (d) showing the presence of gas in the soft tissues and the blood vessels of the brain. (e) and (f) Images obtained during gas sampling showing needles in the sampling position in the cardiac cavities (e) and the pleural cavities (f).
samples collected during the autopsy revealed elevated values of C-reactive protein (175 mg/L), consistent with an inflammatory state and a procalcitonine level compatible with a bacterial infection (0.69 μg/L). The results also revealed signs of cardiac dysfunction with a very high level of N-terminal prohormone of brain natriuretic peptide (>35 000 ng/L) and cardiac necrosis with a very high troponin-T level (>56 000 ng/L).

**Autopsy with immersion of the heart in water (Richter’s technique)**

The autopsy was performed by one board-certified forensic pathologist and one forensic pathologist in training. Opening of the thoracic cavity revealed numerous gas bubbles in the fatty tissue covering the heart. The pericardium was opened and filled with water, and the heart floated. An incision was made in the right ventricle with a scalpel, and blood with gas bubbles escaped. This technique was performed mainly for training purposes and not for gas sampling, as the gas from the cardiac cavities had already been sampled during CT imaging. The autopsy revealed the presence of numerous gas bubbles throughout the vascular system, even in the small vessels of the brain (Figure 2). Additionally, changes to the pulmonary parenchyma were consistent with pneumonia. No other major findings were made.

**Gas analysis**

An Agilent 6890N GC (Agilent Technologies, Palo Alto, CA) combined with a headspace gas autosampler and equipped with an Agilent Select Permanent Gases column arrangement was used. This column arrangement is specially designed for gas analysis and contains a molecular sieve 5 A PLOT capillary column (10 m × 0.32 mm i.d.) and a Porabond Q column (50 m × 0.53 mm i.d.) in parallel, which allows for separation of carbon dioxide (CO₂). The column temperature was maintained at 45 °C for 13 min. The injector temperature was 100 °C, and the injection was conducted in splitless mode. Helium was used as the carrier gas at a constant flow rate of 8 mL/min. The gas detection and quantification were performed with a thermal conductivity detector set at 150 °C. The system was calibrated for each gas with standard gases of H₂S (Multigas, Domdidier, Switzerland), O₂ and N₂ (from laboratory air), and CH₄ and CO₂ (Carbagas, Lausanne, Switzerland). With this system, all the gases could be
detected in the same run. The gas compositions for the different intracadaver sampling sites are displayed in Figure 3.

**Histological analysis**

Samples of the brain, heart, lungs, liver and kidneys were taken during autopsy and stained following a standard haematoxylin and eosin protocol. The heart samples were taken from both ventricles, and the interventricular septum exhibited epicardial vessels deprived of red cells and small intramyocardial haemorrhages. The lung samples taken from each lobe contained numerous clusters of leucocytes, mainly neutrophils, and, to a lesser extent, some macrophages, predominantly in the left lower and right upper lobes. These findings are consistent with the pneumonia suspected on admission to the hospital. There were no significant findings from the other organ samples.

Samples of the heart were prepared and analysed for deposition of the plasma antigen fibronectin and the terminal complement complex C5b-9 to look for early cardiac damage, especially the right ventricular ischemia. None of the samples showed reaction for C5b-9, and only small groups of cells stained for fibronectin. Based on these results, cardiac ischemia was not likely in this case.

**Medico-legal conclusion**

In light of the different results, the cause of death was attributed to a fatal O2 embolism. This was caused by the infusion of pure O2 from the wall into the vascular system through a venous catheter connected to a nasal cannula. Although from a medico-legal point of view the circumstances of the death (accident, suicide and homicide) remained unclear, police investigations led to the suspicion that the patient had connected the

---

**Figure 3.** Results of gas analysis showing the different gas compositions at intracadaver sampling sites. O2: Oxygen; N2: Nitrogen; and CO2: Carbon dioxide.
nasal cannula to the venous catheter. The wound on the forehead and the bruises observed during the external examination could be explained by a fall from or next to the hospital bed, as suggested by the police.

**Discussion**

From the beginning of the medico-legal investigation, especially during the external examination, given the extended subcutaneous emphysema, a massive gas embolism was suspected. This was later confirmed by the results of the post-mortem investigations. The CT scan revealed the presence of massive gas embolism in the cardiac cavities. Gas was also present in the thoracic and abdominal organs and cavities, and in the vascular system, to a similar extent to that which is usually encountered in severely decomposed bodies. Given the short post-mortem interval in this case, an exogenous source of gas, O₂, or air was the most probable explanation of such a large volume of gas in the body.

After the autopsy, cardiac gas embolism was suspected as the cause of death. The organs did not show any sign of advanced putrefaction, which ruled out putrefaction as the sole source of gas in the body. The histological, immunohistochemistry and post-mortem chemistry results excluded cardiac ischemia or severe sepsis as the cause of death. Intoxication as the cause of death was excluded based on the results of toxicological analyses.

Following these exclusions, the only remaining question was the type of the gas responsible for the fatal embolism. In the gluteal tissue, the pectoral muscle, the scrotum and the abdominal cavity, very high concentrations of O₂ (>60% or >25 μmol/mL) were measured. In addition, O₂ concentrations in the carotid artery and jugular vein were between 20% and 30% (8.3–12.5 μmol/mL). These O₂ concentrations are related to the cause of death. However, unexpectedly low O₂ concentrations were found in the thoracic aorta, thoracic cavity and heart, and especially in the two last sites (<20% or <8.3 μmol/mL). Conversely, the CO₂ concentrations were unexpectedly high in these three sampling sites, ranging from 20% to 40% (8.3–16.6 μmol/mL). High concentrations of CO₂ (20%–30% or 8.3–12.5 μmol/mL) have been found in experiments with New Zealand white rabbits euthanized by air embolism [13]. Importantly, similar CO₂ concentrations have been found in scuba diving fatalities [7] and in animals [13,14], and are independent of the gas composition (e.g. air, nitrox, helium).

We proposed the increase in gaseous CO₂ in the heart could be explained by the bicarbonate buffer system, with a huge reservoir of dissolved CO₂ in the blood (Figure 4). Post-mortem acidification of the body caused by decomposition may push the equilibrium of this system towards the conversion of bicarbonates into CO₂. Intravascular active gas release (pure O₂ in this case), a post-mortem off-gassing effect, or arterial gas embolism following barotrauma in scuba divers may flush the blood and carry the gaseous CO₂ throughout the body. In relatively airtight organs such as the heart, the gas could accumulate, whereas it is

![Diagram](image_url)

**Figure 4.** Possible mechanism for the increase of gaseous CO₂ in the heart.
less likely to accumulate in open systems such as veins and arteries. This could explain the high O2 concentrations found in tissues with low blood flow (e.g. the gluteal tissue, the scrotum and the abdominal cavity).

The gas sampling and analyses allowed for the exclusion of decomposition as the origin of the intra-cadaver gas. No changes in gases such as hydrogen, hydrogen sulphide or methane were identified. Consequently, the CT scans can be interpreted by the forensic radiologist/pathologist to support gas embolism as the cause of death. The very high O2 concentrations and CT scan results are consistent with a pure O2 fatal gas embolism [15]. The CO2 levels observed are consistent with the data found in the literature. However, more research should be performed to understand the post-mortem presence of gaseous CO2.

Conclusion

In conclusion, our case report presents a case of a fatal O2 injection. The analysis of gas samples allowed us to confirm the police hypothesis that the person died because of O2 injection. High CO2 levels were observed, and a mechanism was proposed for this phenomenon. This case illustrates how post-mortem gas analyses are very important for medico-legal investigations.

Acknowledgments

We thank Mrs Milena Uehlinger for her precious help in editing this manuscript.

Disclosure statement

No potential conflict of interest was reported by the authors.

Funding

The authors received no funding for this research.

References

[1] Jeffery AJ. The role of computed tomography in adult post-mortem examinations: an overview. Diagn Histopathol. 2010;16:546–551.
[2] Krantz P, Holtas S. Postmortem computed tomography in a diving fatality. J Comput Assist Tomogr. 1985;7:132–134.
[3] Chevallier C, Doenf F, Vaucher P, et al. Postmortem computed tomography angiography vs. conventional autopsy: advantages and inconveniences of each method. Int J Legal Med. 2013;127:981–989.
[4] Grabherr S, Grimm J, Baumann P, et al. Application of contrast media in post-mortem imaging (CT and MRI). Radiol Med. 2015;120:824–834.
[5] Egger C, Bize P, Vaucher P, et al. Distribution of artificial gas on post-mortem multidetector computed tomography (MDCT). Int J Legal Med. 2012;126:3–12.
[6] Egger C, Vaucher P, Doenf F, et al. Development and validation of a postmortem radiological alteration index: the RA-Index. Int J Legal Med. 2013;225:53–59.
[7] Varlet V, Smith F, de Froidmont S, et al. Innovative method for carbon dioxide determination in human postmortem cardiac gas samples using headspace-gas chromatography–mass spectrometry and stable labeled isotope as internal standard. Anal Chim Acta. 2013;784:42–46.
[8] Varlet V, Smith F, Giuliani N, et al. When gas analysis assists with postmortem imaging to diagnose causes of death. Forensic Sci Int. 2015;251:1–10.
[9] Varlet V, Bruguier C, Grabherr S, et al. Gas analysis of exhumed cadavers buried for 30 years: a case report about long time alteration. Int J Legal Med. 2014;128:719–724.
[10] Lauer E, Villa M, Jotterand M, et al. Imaging mass spectrometry of elements in forensic cases by LA-ICP-MS. Int J Legal. 2017;131:497–500.
[11] Grabherr S, Doenf F, Steger B, et al. Multi-phase post-mortem CT angiography: development of a standardized protocol. Int J Legal Med. 2011;125:791–802.
[12] Schneider B, Chevallier C, Dominguez A, et al. The forensic radiographer: a new member in the medico-legal team. Am J Forensic Med Pathol. 2012;33:30–36.
[13] Bernaldo de Quirós Y, González-Díaz O, Møllerløkken A, et al. Differentiation at autopsy between in vivo gas embolism and putrefaction using gas composition analysis. Int J Legal Med. 2013;127:437–445.
[14] Pierucci G, Gherson G. Studio sperimentale sull’embo-lia gassosa con particolare riguardo alla differenzia-zione fra gas embolico e gas putrefattivo [Experimental study on gas embolism with special reference to the differentiation between embolic gas and putrefaction gas]. Zacchia. 1968;44:347–373.
[15] Laurent PE, Coulange M, Desfeux J, et al. Post-mortem computed tomography in a case of suicide by air embolism. Diagn Interv Imaging.2013;94:460–462.