Abstract: Toluene and xylene are aromatic hydrocarbons commonly used as an industrial solvent for the manufacturing of pharmaceuticals, paints, and chemicals. The Occupational Safety and Health Administration has determined that toluene levels of 2000 parts per million (ppm) are considered dangerous to life and health. Several studies have examined the absorption of toluene and xylene following inhalation and oral ingestion in humans. Volatile organic compounds that are absorbed into the blood are distributed throughout the body; in particular, distribution of absorbed toluene and xylene in humans and rodents is characterized by preferential uptake in well-perfused and lipophil tissues such as the brain, liver, lungs, and body fat and also in central nervous system. The available studies indicate that xylenes are rapidly absorbed independently from the kind of exposition. We illustrate a fatal case of self-poisoning by ingestion of varnishes diluting solvents, reporting the identification and quantification of volatile organic compounds (toluene, o-m-p xylene) from human biologic liquids and viscera samples using the Solid-Phase Microextraction-Headspace-Gas Chromatography/Mass Spectrometry to toxicological analysis, and the histopathological findings evaluated in liver, kidney, and lungs.

Key Words: toluene, xylene, paint thinner ingestion, self poisoning, varnish-diluting solvents

CASE REPORT

A Fatal Case of a Paint Thinner Ingestion

Comparison Between Toxicological and Histological Findings

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Toluene and xylene are aromatic hydrocarbons (C₇H₈ and C₈H₁₀) commonly used as an industrial solvent for the manufacturing of pharmaceuticals, paints, and chemicals. The Occupational Safety and Health Administration has determined the acceptable level of occupational exposure to toluene and xylene for people in the workplace; toluene levels of 2000 ppm are considered dangerous to life and health. Several studies have examined the absorption of toluene and xylene following inhalation and oral ingestion in humans; volatile organic compounds (VOC’s) that are absorbed into the blood are distributed throughout the body.

Distribution of absorbed toluene and xylene in humans and rodents is characterized by preferential uptake in well-perfused and lipophil tissues such as the brain, liver, lungs, and body fat. Data from literature suggest that, after oral ingestion, toluene accumulates in the liver, whereas after inhalation toluene accumulates in the brain. Studies in humans and animals show that the central nervous system is also the main target of xylenes. The various isomers o-, m-, p-xylene have similar health effects.

The available studies indicate that xylenes are rapidly absorbed independently from the kind of exposition; once absorbed the toxins undergo a considerable metabolism, being the liver the primary site of such metabolism and hippuric acid the main metabolite. As concerns quantification of VOCs in postmortem matrices several methods have been proposed. These substances have been generally determined by gas chromatography after extracting the compounds using solvent extraction methods or even static and dynamic headspace (HS) techniques. The headspace techniques include HS-solid phase microextraction (HS-SPME), cryogenic oven trapping (HS-COT), cryogenic focusing (HS-CF), and purge and trap. SPME has proved to be an excellent sampling technique for analysis of volatile organic compounds. In fact, SPME is solvent-free and require only minimal sample handling and preparation. These technique uses polymer coated/fused-silica fibers for the extraction of organic compounds from different matrices.

CASE REPORT

A 15-year-old white boy was found in supine position in the garden of his Teacher’s country-house. The student was immediately taken to the local emergency room and, after a few minutes, he was pronounced dead.

History indicated that the boy, after failing an examination, had gone to his teacher, to talk to him. After a few minutes, the boy fell on his back, with agonal gurgling and foam in his mouth. After his death, the local police-officers found numerous small bottles of varnish-diluting solvents.

By an order of the legal authorities, the external examination and the autopsy were performed 2 days later at the Institute of Legal Medicine of Palermo. External examination: the young boy was 175 cm tall and his weight was 65 kg. No injuries were found in his body; only a nasal hemorrhage and labial and subungual cyanosis were found.

MATERIALS AND METHODS

Autopsy Findings

The forensic autopsy revealed cerebral edema and congestion of cerebral veins. There were no lesions in the scalp or the galea capitis and no intracerebral hemorrhage. Pulmonary edema and pancreatic and renal congestion were found. The gastric content was brownish liquid (300 mL) and had an odor of organic solvent. Nothing else was found during the autopsy. The blood alcohol screening was negative. Urine samples were too exiguous to perform any analytical screening.

Histologic Findings

Samples of organs taken during autopsy were fixed in buffered formalin 10% for a period of 2 weeks and then processed. From blocks obtained, sections thick 4 to 6 μm were cut and colored with hematoxylin-eosin. The microscopic examination showed severe cerebral...
edema, diffuse axonal damage, diffuse cytotoxic edema, and multifocal ischemic neuronal injury. Renal parenchyma showed cellular swelling and tubular necrosis (Fig. 1). Hepatic parenchyma showed diffuse vacuolar degeneration of hepatocytes and vascular congestion (Fig. 2).

Diffuse vascular congestion, intra-alveolar hemorrhage, and edema were observed in pulmonary slides (Fig. 3). Most part of red blood cells was lysed and alveolar septa were fragmented probably due to direct solvent activity on cellular membranes. The microscopic examination of esophagus showed multifocal erosion of epithelium and severe vascular congestion and edema of mucous and submucous membranes.

**Toxicological Findings**

A Supelco (Bellefonte) SPME manual fiber assembly with fused-silica (85 μm film thickness) carboxen-polydimethylsiloxane-coated (PDMS) was used for all analysis.

The standards of toluene, toluene D₈ (99.96%) and o-, m-, p-xylene (99.8%) were obtained from Sigma-Aldrich (Germany). All of the reagents used in this study were of HPLC grade. Water and acetonitrile were bought from Fluka (Germany).

**Determination and quantitation of volatile substances was performed by HS/Solid-Phase Microextraction/Gas Chromatography-Mass Spectrometry (HS-SPME-GC/MS).**

Analyses were carried out with GC/MS equipped with a column Supelcowax 10 TM (30 m × 0.25 mm ID, 0.25 μm film thickness-Supelco).

**Headspace SPME Sampling Procedure**

All of the results of postmortem samples were stored at –20°C up to the analysis. Extraction of volatiles was first optimized by choice of fiber. Several studies showed that Carboxen/PDMS coated fibers have been employed for biomonitoring of benzene and toluene in human blood. However in the preliminary studies, extractive capacity of Carboxen/PDMS (CAR/PDMS) and PDMS fibers was evaluated, and CAR/PDMS gave the highest level of these substances.

The equilibration and adsorption times were established after several tests on samples of blood, liver, lung, kidneys, and gastric content (Table 1).

Based on the preliminary experiments, all of samples up to the analysis were kept at a temperature of 40°C ± 0.5°C in aluminum block-heater for 1 hour.

**Instrument Conditions and GC-MS Analysis**

A preliminary study on gastric content as well as toxic determination in gastric content and blood were conducted using a single quadrupole instrument (HP 5890-Hewlett Packard).

Afterward, the analyses on the other tissue samples (liver, kidney, and lungs) were performed using a Varian STAR 3400 CX GC coupled with an ion trap MS analyzer (Varian Saturn 3 GC-MS).

<table>
<thead>
<tr>
<th>Matrices</th>
<th>Equilibration Time</th>
<th>Adsorption Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood</td>
<td>10 min</td>
<td>2 min</td>
</tr>
<tr>
<td>Gastric content</td>
<td>30 min</td>
<td>60 min</td>
</tr>
<tr>
<td>Liver</td>
<td>30 min</td>
<td>60 min</td>
</tr>
<tr>
<td>Kidney</td>
<td>30 min</td>
<td>30 min</td>
</tr>
<tr>
<td>Lung</td>
<td>30 min</td>
<td>60 min</td>
</tr>
</tbody>
</table>

**Figure 1.** (EE 200×) Renal parenchyma with diffuse acute tubular necrosis.

**Figure 2.** (EE 400×) Hepatic parenchyma showing diffuse vacuolar degeneration of hepatocytes and sinusoidal congestion.

**Figure 3.** (EE 200×) Pulmonary parenchyma with alveolar edema and acute alveolar hemorrhage.
and an autosampler (Varian 8200 CX), to achieve a better reproducibility on data and reduce manual errors.

The SPME fiber was placed into a injector (splitless mode) and the injection temperature was 260°C. Carrier gas flow was 35 cm/s, column pressure was 56 KPa, the oven program condition were the following: isotherm 50°C for 12 minutes, ramp up to 250°C at 40°C/min, isotherm at 250°C for 3 minutes.

The EI/MS spectra (70 eV) were carried out operating in SIM mode and as extracted ions, using respectively a single quadrupole instrument and ITD mass analyzer. In both instruments, full scan mode was used as an identification test. Scan time was fixed on 1 second Monitored ions were m/z 78 (benzene); m/z 98 (toluene D₈); m/z 91 (toluene); and m/z 91, m/z 106 (xylenes).

To assess the reproducibility of the analysis, each sample was analyzed 3 times.

The preliminary analysis allowed an estimate of the amount of the various analytes in the different matrices, in this way, it was possible to determine the amount of standard addition.

Liquid samples and particularly blood and gastric fluid were analyzed using the multiple standard addition method, to reduce to the minimum the matrix effect. For the solid samples, frozen kidney and liver quantification was performed using a calibration curve benzeno D₈ as internal standard.

The general approach was to optimize the method to avoid as possible analyte losses, minimizing sample manipulation. This required a different approach, depending on the tissues analyzed to obtain a best accuracy of measure. However, similar chromatographic results were obtained for all the samples and both quadrupole and ITD/MS. A typical example is reported in Figure 4.

**Gastric Content**

Four identical vials containing 455 μL of samples were prepared. In each one an aliquot of 20 μL of internal standard (Toluene D₈ at concentration 2000 ppm) was added. Further, an appropriate amount of standard solution was added to obtain a final standard concentration of 500, 1000, and 2000 ppm of toluene; 150, 300, and 600 ppm of ortho- and meta-xylene; and 50, 100, and 200 ppm of para-xylene, respectively, in the final volume of 600 μL. This was obtained by further diluting the sample with HPLC grade water.

**Blood**

Four vials (2 mL) containing 0.5 g of sample were prepared. To each one an aliquot of 5 μL of internal standard (benzene at concentration 200 ppm) was added.

An appropriate amount of standard solution was added to obtain a final standard concentration of 5, 10, 15 ppb of toluene; ortho-, meta-, and para-xylene in the final volume of 1 g. This was obtained by further diluting the sample with HPLC grade water.

**Liver, Kidney, and Lung**

In this case, sample handling was particularly difficult due to the volatility of the analytes and the intrinsic nature of the matrix. For such solid samples, a homogenization process was necessary, which was conducted in ice bath to avoid significant volatile losses. Even cautiously homogenized, the samples were difficult to manage. It was particularly difficult to weight small amounts necessary for autosampler vials and perform for each sample a multiple standard addition. Several attempts were made but the method adopted lacked of precision, results obtained were used as an estimate of the levels of toxic in the liver, kidney, and lung tissues. A new strategy for these samples was then adopted and quantification was made through and external calibration curve. To minimize the errors due to matrix effect, the calibration curve was generated trough a spiked simulated.

Matrix made ad hoc for each tissue analyzed. Thus, homogenized porcine liver, bovine lung, and kidney were used as matrices. Further, for better accuracy, toluene D₈ was adopted, as internal standard. For lung, the preliminary study confirmed that concentration levels were well below the quantification level and for several analytes no analytical signal was detected. Instead for the liver and kidney, this study revealed as raw estimate the following concentrations reported below.

The animal samples (liver and kidney) have been frozen at −23°C for 1 week (with the aim to better simulate postmortem samples) and afterward have been homogenized in ice bath.

A standard mix (A) solution containing toluene at about 1000 ppm (100 μL in 100 mL water/acetonitrile 60:40), ortho- and meta-xylene at 330 ppm (33 μL in 100 mL water/acetonitrile 60:40) and para-xylene at 100 ppm (10 μL in 100 mL water/acetonitrile 60:40) was prepared to be added to liver samples.

Another standard mix (B) solution containing toluene at about 25 ppm (25 μL in 100 mL water/acetonitrile 60:40), para-xylene at about 25 ppm (25 μL in 100 mL water/acetonitrile 60:40), meta-xylene and ortho-xylene at about 100 ppm (100 μL in 100 mL water/acetonitrile 60:40) was prepared to be added to kidney samples. As it can be seen, the standard solutions have been prepared volumetrically, therefore, the following concentrations reported in ppm are corrected for toluene, ortho-, meta-, and para xylene density.

For the calibration curve, aliquots of 600 mg of homogenate were prepared and, in each sample, an appropriate aliquot of standard mix solution (A for liver and B for kidney) (Table 2) was added.
The samples were further diluted to 30 mL (for liver) and to 10 mL (for kidney) using HPLC grade water, and vortex mixed for 1 minute. Vials containing 200 μL of diluted samples added with 20 μL of internal standard solution (toluene-D₈ 2000 ppm in water/CH₃CN 60:40) were finally used for analysis.

Calibration curves obtained reporting concentration versus standard/internal standard area ratio are shown in Figure 5. Calibration levels were optimized to achieve at least one point of the calibration curve in proximity of the expected concentration for each toxic in the tissue, few concentration levels per curve were taken into consideration as it was known the order of magnitude of analytes. Each measure was repeated twice to check the reproducibility.

Human samples were homogenized and 600 mg of tissue homogenate was diluted to 30 mL (for liver) and to 10 mL (for kidney) using HPLC water; vials were prepared using 200 μL of diluted sample added with 20 μL of internal standard solution (toluene-D₈ 2000 ppm in water/CH₃CN 60:40). Results are reported in Table 3.

### DISCUSSION

Organic compounds are distributed throughout analyzed tissues reflecting the greatest affinity for lipid-rich tissues such as liver and kidney instead that for blood (Table 2).

Gastric content showed very high concentration of toxics, even if a relevant part of ingested solvents were also distributed in body tissues. The highest concentration of solvents was in liver according to lipophilic nature of analyzed parenchyma.

Within kidney toluene concentration was lower than xylenes concentration.

Only a few concentration (<LOQ) of organic compounds were detected in lung, probably because of reanimation, time passed between autopsy and ingestion, and also the wide exchange surface of pulmonary tissue.

Quite surprisingly, just in blood (Fig. 6), the presence of further aromatic compound was exploited. This was identified as...
TABLE 3. VOCs Concentration in the Various Tissues in ppm

<table>
<thead>
<tr>
<th>Sample</th>
<th>Toluene ppm</th>
<th>Ortho-Xylene ppm</th>
<th>Meta-Xylene ppm</th>
<th>Para-Xylene ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastric content</td>
<td>21,500</td>
<td>1500</td>
<td>1075</td>
<td>5400</td>
</tr>
<tr>
<td>Blood</td>
<td>0.060</td>
<td>0.232</td>
<td>0.160</td>
<td>0.065</td>
</tr>
<tr>
<td>Liver</td>
<td>41.5</td>
<td>12</td>
<td>15</td>
<td>6</td>
</tr>
<tr>
<td>Kidney</td>
<td>0.939</td>
<td>4.15</td>
<td>3.86</td>
<td>1.08</td>
</tr>
<tr>
<td>Lung</td>
<td>&lt;LOQ</td>
<td>&lt;LOQ</td>
<td>&lt;LOQ</td>
<td>&lt;LOQ</td>
</tr>
</tbody>
</table>

FIGURE 6. Chromatogram of a blood sample spiked with: 1 benzene, 3 toluene, 4 p-xylene, 5 m-xylene, 6 o-xylene.

ethylbenzene by mass spectrum and research in Nist 02 library. The presence of ethylbenzene was revealed only in blood, none of the other tissues under investigation revealed traces of this organic compound. We, up to now, have not explanation for the presence of this further contaminant since all standard have been checked to verify its presence even as trace contaminant. Ethylbenzene was found neither in liver and kidney and, most surprisingly, neither in gastric content, were concentration of the solvents was much higher.

Because of their lipophilic character, volatiles have a serious impact in the brain and in other parts of the nervous system. They act as a central nervous system depressant and this is the most common cause of death in VOC’s ingestion. The lipid solubility, volatility, and route of exposure of the compounds enhance their toxicity.16 The most important toxicological effects are observed for the heart, brain and neurologic system, liver, kidney, and lungs.17 Little is known about the mechanisms by which toluene produces acute effects but it is reasonable to assume that its toxic effects are due, at least in part, to its general characteristics as a solvent: the presence of solvent molecules in cholesterol-filled interstices between phospholipids and sphingolipids changes membrane fluidity, thereby altering intercellular communication and normal ion movements.18

An alternative hypothesis is that toluene partitions into hydrophobic regions of proteins and interacts with them, thereby altering membrane-bound enzyme activity and/or receptor specificity.19

Reports of oral exposure to toluene in humans are limited to case reports of accidental acute ingestion. In humans, a toluene ingestion (only few mL) is fatal and death occurs after only 30 minutes.20 A case report of a 51-year-old man who died approximately 30 minutes after he had ingested a large quantity of toluene (625 mg/kg) was presented; the probable cause of death was severe central nervous system depression, they also noted acute tubular necrosis and acidosis.21 Caravati and Bjerk (1997)20 report on a nonlethal case of a 46-year-old man who had ingested approximately one-quarter of paint thinner containing toluene. The patient presented a severe central nervous system depression, severe abdominal pain, diarrheal, hemorrhaghe gastritis, acute tubular necrosis, and acidosis. Human studies following oral exposure to xlenes are not available.

Abu-Al-Ragheb et al (1986)22 report on a 27-year-old man who committed suicide by ingesting xylenes. Histopathological findings included areas of pulmonary edema and congestion. The probable cause of death was attributed to respiratory failure and asphyxia, a secondary response elicited by depression in the respiratory center in the brain.

In another case, accidental ingestion of xylenes resulted in a deep coma lasting more than 26 hours, hepatic impairment, hematemesis, acute pulmonary edema, and other pulmonary complications.23 Our case showed multiorgan damage and diffuse cytotoxic edema. The probable cause of death was attributed to acute toxic effect as solvent in body, and to pulmonary complications, respiratory failure, depression in the respiratory center, and extensive cerebral edema.

CONCLUSIONS

The case study finding provides additional support to the proposal that fatal intoxication by ingestion of aromatic solvents can occur in a subject having suicidal intent.

Moreover, the comparison of histologic and toxicological findings provides to understand correlation between organic damage following ingestion and amounts of solvents found. Findings in toxic concentration in various tissues indicate that the distribution of toxic is related to their lipophilicity. However, in lung concentration of solvents was particularly low and far below to the limit of quantization of the method. This imply a small partition of the toxic in lung otherwise that lung large exchange surface or the reanimation processes or both of them could be responsible of the elimination of these solvents in this tissue.

REFERENCES

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