Elimination of toluene from venous blood and adipose tissue after occupational exposure

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ABSTRACT In a group of 37 rotogravure printers a close correlation (r=0.78) was found between the time weighted toluene exposure during a five day working week (range 8–416 mg/m³, median 75) and the concentration of toluene in subcutaneous adipose tissue (range 1·1–20·7 mg/kg, median 3·8). After exposure ceased, the elimination of toluene was followed up in 11 subjects. The toluene concentration in venous blood decreased non-linearly and the elimination curves contained at least three exponential components. The first two had median estimated half times of nine minutes and two hours respectively. The third component, with a median half time of 90 hours, reflected the decline in adipose tissue, which had a median half time of 79 hours (range 44–178). The study showed protracted endogenous toluene exposure from adipose tissue deposits long after the end of exogenous exposure. The observations also suggest that the blood toluene concentrations on Monday mornings might be used as an index of the exposure in the previous week.

A knowledge of the kinetics for eliminating toluene from venous blood and adipose tissue is important in evaluating toxicity and for planning biological monitoring. The concentration of toluene in venous blood (B-toluene) is used for the biological monitoring of occupationally exposed workers.1,2 The elimination of toluene from arterial and venous blood has been investigated in experimentally exposed subjects, showing a rapid decrease of toluene in both arterial and venous blood.3–5 After 60 minutes, only 9% was left in alveolar blood and 23% in venous blood.4 The elimination of toluene from subcutaneous adipose tissue has also been studied in voluntary subjects exposed under experimentally controlled conditions. The half time for toluene in subcutaneous adipose tissue ranged between 0·5 and 2·7 days (12–64·8 h).6

Little is known, however, about the elimination of toluene in occupationally exposed subjects. In a previous study of plastic processing workers an accumulation of toluene in the blood during a working week was found at an exposure level of 184–332 ppm (700–1270 mg/m³).7 Brugnone et al suggest an accumulation at a mean exposure level of 146 mg/m³.1 We have previously shown an accumulation of toluene during a working week at exposure levels well below the current Swedish TLV (300 mg/m³), which suggests a long half life.2

In the present investigation the association between toluene in environmental air and subcutaneous adipose tissue was studied in occupationally exposed subjects. Toluene elimination was followed up in blood during a weekend and in blood and adipose tissue during a week’s holiday.

Material and methods

SAMPLING STRATEGY

Sampling 1—Environmental air was monitored during a five day working week in two rotogravure printing factories. Samples of subcutaneous adipose tissue were taken from the workers towards the end of the week. Biopsy specimens were taken directly after work (25 on Thursday, 14 on Friday).

Sampling 2—To investigate elimination 11 rotogravure printers voluntarily took part in samplings 2 and 3. Fast elimination from venous blood was studied during a weekend. Blood was sampled directly after work and then after 10, 20, 40, 60, 120, 180, 240, and 360 minutes. Two further samples were taken 21 and 72 hours after exposure ceased.

Sampling 3—Elimination from adipose tissue and the slow decline in venous blood was studied during a week’s holiday. Sampling was performed directly after work and then after 63 and 135 hours without exposure. For the 11 participating subjects the amount of body fat was also determined by means of skeletal...
measurements according to the method described by von Döbeln.89

SUBJECTS
The exposed subjects in sampling 1 consisted of 37 male printers exposed to toluene for 2–43 years (median 23). They were aged from 22 to 62 (median 45). In samplings 2 and 3 the group examined consisted of 11 male printers exposed for 13–36 years (median 24). They were aged from 30 to 62 (median 47).

The printers’ state of health, use of drugs, smoking habits, and alcohol consumption were checked with a medical examination and a detailed interview.

VENOUS BLOOD SAMPLES
Blood was sampled in heparinised tubes. The samples were analysed with the head-space technique, using ethylbenzene as an internal standard, on a gas chromatograph equipped with a flame ionisation detector (FID) (Varian model 3700). Standards were made from venous blood to which a known amount of toluene had been added. The detection limit was 0.01 μmol/l. The precision of the method was 4%, both at B-toluene of 0.20 and 3.8 μmol/l.

SUBCUTANEOUS ADIPOSE TISSUE SAMPLES
Skin was penetrated by a scalpel without use of local anaesthetic in the upper lateral gluteal region. Subsequently, biopsy specimens were obtained by a lance shaped needle with dimensions 2.0 × 80 mm (KN 1480 W, Terumo Europe NV, 3030 Leuven, Belgium). The needle was connected via a luer lock to a 50 ml syringe with a glass barrel and plunger (B-D Yale, Becton, Dickinson & Co).

The concentration of toluene in adipose tissue was determined by gas chromatography after evaporation of the solvent at 150°C into nitrogen (1 ml/min) which was continuously exchanged.10 The evaporation lasted 30 minutes and the gas was collected into 30 ml all glass syringes. The gas samples were analysed by a gas chromatograph (Carlo Erba model 4160) equipped with a quartz capillary column DB-1701 (length 30 m, inner diameter 0.25 mm). The toluene concentrations were calculated on the basis of standard air samples with known toluene concentrations. The error of the method for single determinations was found to be within 11% of the mean value of ten duplicate determinations with different puncture channels and the concentrations ranging from 0.5 to 10.2 mg/kg. To obtain reliable results, blood contaminated specimens or those weighing less than 10 mg, or both, were excluded. The mean loss of weight during evaporation was 8% (range 1–18%).

The specificity of the analysis of toluene in body fat has been checked in two ways. Firstly, small amounts of pure toluene were added to the nitrogen after evaporation of the tissue. No difference in retention time was found for the evaporated and added toluene. Secondly, biopsy specimens from one person contributing the exposure measurements were taken before and after visiting a toluene contaminated area. Toluene was only found in samples taken after visiting the plant.

ENVIRONMENTAL AIR SAMPLES
The environmental concentration of toluene was estimated by personal sampling (sampling 1). These workers wore a personal sampler with which samples were continuously taken from the ambient air at 30 minute intervals (the Linder Gaspirator, Instrument AB Lambda, Sweden). The sampler sucked air from the breathing zone through a Teflon capillary tube into a 30 ml all glass syringe.11 The toluene content of the air was immediately determined with a portable gas chromatograph equipped with FID (AID, model 511). Standard gases were prepared by adding known amounts of toluene into a glass bottle with a known volume of clean air. The error of the method, calculated from 25 double determinations ranging from 67 to 335 mg of toluene/m³, was within 1-1% of the mean value.

Data analysis

The decreasing levels of B-toluene in each subject were fitted as a three compartment exponential model, that is, a linear combination of three exponentials with different rate coefficients (and thus half times). The non-linear regression procedure of the BMDP package (P3R) was used. It uses the iterative Gauss-Newton algorithm.12 Owing to the relatively short follow up (sampling 2), the half time for the slowest compartment was estimated separately, using data collected over an independent and longer period (sampling 3) and held fixed in the three compartment model. Two parameters were estimated for each compartment: an elimination rate, transformed and quoted as the half time tj (1), tj (2), and tj (3) and the amount corresponding to each compartment Y (1), Y (2), and Y (3).

Associations were investigated with Spearman’s rank correlation coefficient. All p values are one tailed.

Results

AIR LEVELS
In all we have exposure data for seven weeks. The ambient air level ranged from 8 to 416 mg/m³ (median 75). For the 11 subjects voluntarily taking part in samplings 2 and 3 the time weighted exposure for this week, based entirely on measured exposure levels, ranged from 35 to 246 mg/m³ (median 115).

TOLUENE IN VENOUS BLOOD
B-toluene was also measured in 21 unexposed work...
Elimination of toluene from venous blood and adipose tissue after occupational exposure

All nine subjects had detectable levels of toluene in venous blood; all had concentrations below our detection limit (0.01 μmol/l). The blood in six of those who smoked, however, contained a trace of a compound with the same retention time in the chromatogram as toluene.

At the end of the exposure after the last shift on the Friday, the median B-toluene in sampling 2 was 2.3 μmol/l (range 1.4–4.4; fig 1). In sampling 3 the corresponding value was 1.0 μmol/l (range 0.22–1.9; fig 1).

The decrease of toluene was non-linear. Furthermore, the decay did not display a simple exponential pattern. After 63 hours (sampling 3), the median B-toluene was 0.12 μmol/l (range 0.03–0.59). After 135 hours toluene was still present in blood (0.06 μmol/l, range 0.02–0.17). In sampling 2 the median B-toluene after 21 hours was 0.30 μmol/l (range 0.18–0.56) and after 72 hours 0.16 μmol/l (range 0.05–0.24).

The median half time for the slow elimination (t1/2 (3)), calculated from the blood samples at 63 and 135 hours in sampling 3, was 90 hours (range 50–324). These individual estimates of the slow elimination of toluene from blood were used for calculating the faster decline. A model with three exponentials showed a better fit than one with two. Convergence was achieved after between 13 and 22 iterations for all subjects. The results of the kinetic calculations for the elimination of toluene in blood are listed in the table. The fastest component had a median estimated half time of nine minutes. From the estimates of the intercept parameter (Y (1)), this compartment constituted a median of 35% (range 20–70) of the maximum B-toluene at the end of exposure. The median half time of the intermediate component was two hours and accounted for the elimination of a median of 42% (range 22–63) of the toluene. The slow component corresponded to a median of 17% (range 3–26).

TOLUENE CONCENTRATIONS IN SUBCUTANEOUS ADIPOSE TISSUE BIOPSY SPECIMENS

At the end of the exposure, the toluene had declined in adipose tissue biopsy specimens (fig 1). In one subject the fat toluene concentration had increased between

<table>
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<th>Subject No</th>
<th>t1/2 (1) (min)</th>
<th>Y (1) (μmol/l)</th>
<th>t1/2 (2) (h)</th>
<th>Y (2) (μmol/l)</th>
<th>t1/2 (3) (h)</th>
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Fig 1  Elimination of toluene: concentrations (medians and ranges) from venous blood (circles) and subcutaneous adipose tissue (squares) after end of exposure.

Friday and Monday. He denied exposure during the weekend. We could not explain the increase by analytical error, thus there had probably been an exchange of his samples. Accordingly, they were excluded from further calculations. The median of the estimated half times for eliminating toluene from adipose tissue was 79 hours for the remaining 10 subjects (range 44–177).

The toluene concentration in adipose tissue taken directly after work on the Thursday and Friday (sampling 1) was associated with the workers’ mean exposure in the previous week (fig 2). Spearman’s correlation coefficient was \( r_s = -0.780 \), \( p < 0.00001 \) (n = 37). The subject marked with an asterisk in fig 2 had a high exposure the day before sampling, but the exposure level could not alone explain his high concentration in both venous blood and adipose tissue.

ASSOCIATION BETWEEN FAT AND BLOOD TOLUENE CONCENTRATIONS

Comparing elimination of toluene from adipose tissue with the decrease corresponding to the slow component.
ent in blood (median half times 79 v 90) indicates that the B-toluene after about 70 hours exposure free time reflects the elimination from adipose tissue (fig 1). The association between the slow half times estimated in blood (t₁/₂ (3)) and the estimated half times for the toluene elimination from adipose tissue is shown in fig 3 (rₛ = 0.46, p = 0.09; n = 10). For six of the ten subjects, the t₁/₂ (3) in blood are within two standard deviations of the half time of toluene in adipose tissue.

There is a weak positive association between the estimated proportion of toluene in blood corresponding to the slow elimination and the concentration of toluene in adipose tissue (70 hours after work in sampling 3) multiplied by the amount of body fat, expressed as percentage of body weight (rₛ = 0.47, p = 0.08). Using instead the B-toluene 70 hours after exposure strengthened the association (fig 4; rₛ = 0.82, p = 0.007).

The difference between B-toluene directly after work in samplings 2 and 3 is probably due to the approaching holiday at sampling 3. The printers reported that they had lower exposure during their last hours of work just before their holiday. Consequently, B-toluene had already started to decrease during the last working hours.

An important finding in the study is the presence of a slow component in venous blood, which had a median half time of 90 hours. This is in agreement with our previous estimates of 72 hours (range 32–92). The slow compartment corresponded to a median of 14% of the total B-toluene at the end of the exposure. Indeed, after 21 hours we measured 13% of the initial B-toluene, which is also in agreement with our previous findings and is also in reasonable agreement with the 7% found by Brugone et al. Hence, during a weekend the slow compartment should dominate the B-toluene concentration already after 24 hours.

The median estimated half time for elimination of toluene from adipose tissue was 79 hours. This is in
good agreement with the findings in an experimental study by Carlsson and co-workers, in which the subjects were exposed for only two hours but at higher air levels. We found an association between the slow elimination (t1/2) from blood and the half-time for the decrease in subcutaneous fat. The elimination from blood, however, seems to be somewhat slower than that from subcutaneous fat. The discrepancy might be due either to an overestimate of the half time in blood because of a not recorded rapid component or to a decay of the biotransformation of toluene towards the end of the study period. After 70 hours without exposure, we found a significant association between the concentration of toluene in subcutaneous fat and in blood. Thus we have strong evidence that the easily monitored slow elimination from blood reflects the disappearance of toluene from adipose tissue.

The prolonged presence of toluene in blood means that there is an endogenous exposure from the adipose tissue depots, which continues for a long time after the end of a workshift. Thus the nervous system is subjected to the pharmacological effects of toluene during nights, weekends, and even during the first part of holidays. One may speculate whether this prolonged endogenous exposure is important in the development of chronic toxic effects.

The concentration of toluene in subcutaneous adipose tissue towards the end of the working week reflects the exposure during the preceding week. Some values appeared to be outliers (fig 2). The explanation may be that an occasional high exposure during the sampling day or the day before, or both, gives an increase of toluene in body fat, whereas the time weighted exposure for the week is affected to a smaller extent. Despite the close correlation between the concentration of toluene in fat and the air level, the fat biopsy method will not be commonly used for biological monitoring, as sampling is too traumatic for routine use. Instead, preshift B-toluene in Mondays might be used as a biological index for the exposure during the previous week. This suggestion, however, has not yet been tested.

The large interindividual variation in the elimination rates from blood is an important observation. It may be partly due to the lapse of time between end of exposure and sampling but is probably also due to genetic or life style factors, or both. The oxidative metabolism of some xenobiotics is known to vary substantially between individuals. Smokers eliminate toluene faster than non-smokers. In the present study, however, the four current smokers had half times evenly distributed within the group range. Alcohol intake during exposure is known to influence the distribution and elimination of toluene. Our subjects were asked not to drink alcohol during the sampling period and traces of ethanol were found in only a few samples.

The varying half times might influence the possibility of evaluating exposure through biological samples. Thus there is a risk of overestimating the environmental exposure for a subject with a long half time. Protracted endogenous exposure may still mean an increased risk for developing adverse effects from the exposure to toluene. Thus the biological sample may be a good index of risk.

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References


12 BMDP statistical software. Los Angeles: Department of Biometrics, University of California, 1981.


