Quantitative relationship between atmospheric phenol vapour and phenol in the urine of workers in Bakelite factories

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Ohtsuji, H., and Ikeda, M. (1972). Brit. J. Industr. Med., 29, 70-73. Quantitative relationship between atmospheric phenol vapour and phenol in the urine of workers in Bakelite factories. A survey in Bakelite factories revealed that the urinary level of total phenol, free plus conjugated, increased in proportion to the phenol concentration to which the subject was exposed in his working environment. This indicates that analysis of urine for phenol is valuable for the monitoring of exposure to phenol in air.

The results also suggest that the human lung takes up phenol very efficiently. The increase in urinary phenol was entirely attributable to conjugated phenol; regardless of exposure, no significant change was observed in free phenol concentration. Ethereal sulphate or ethereal glucuronide had poor validity as an index of exposure.

Since the beginning of this century the use of synthetic resins has been spreading in every field of life. Bakelite, or phenol resin, is among the most popular resins, even though somewhat old-fashioned.

Following previous reports on the metabolism of styrene (Ohtsuji and Ikeda, 1971) and the use of analyses of urinary metabolites for monitoring the exposure of workers in styrene resin factories (Ohtsuji and Ikeda, 1970), we report, in the present paper, that the urinary concentration of phenol (i.e., total phenol or, preferably, conjugated phenol) is a good index of exposure to atmospheric phenol vapour in Bakelite factories.

Materials and methods

Urine samples
The survey was conducted in the autumn. Urine samples were collected from male workers who had been engaged in laying together paper sheets impregnated with phenol resin to produce Bakelite boards. The workers were exposed to phenol from 8.00 a.m. to noon and from 1.00 p.m. to 4.00 p.m. daily. This type of workshop was chosen for study as the labour was performed in closed rooms where environmental concentrations of phenol were fairly stable. Samples from office workers were also collected for examination of levels of metabolites in controls.

Analysis of urine Phenols (total, free, and conjugated) were determined colorimetrically by the method previously described (Ikeda, 1964) using Gibbs' colour reagent. The specificity of the method and the origin of phenol in urine have been discussed by van Haaften and Sie (1965). The method does not record p-cresol.

Ethereal glucuronide was measured as previously described (Ikeda, 1964) with naphthoresorcinol as a colour-developing reagent.

Determination of ethereal sulphate was based on the method of Bertolacini and Barney (1957) as modified by Ohmori and Hara (1966). Inorganic sulphate was determined on 0-15 ml of urine, which was mixed in a tube with 0-45 ml of water, 0-6 ml of 1 M acetate buffer (pH 4-0), and 0-3 ml of 1 mg/ml potassium oxalate to remove interfering calcium ion. After the addition of 5-4 ml of 31-7% ethanol/water (v/v) and about 20 mg of barium chloranilate, the tube was shaken vigorously for 15 minutes and centrifuged. The red colour was measured at 530 nm.

For the determination of total sulphate, urine was sealed in an ampoule with 3 volumes of 1/3 N HCl, and heated at 100° C for 1 hr. A portion, 0-6 ml, of the hydrolysate was added to 1 M acetate buffer (pH 4-0) and enough 0-25 N NaOH to bring the pH to 4-0. Sulphate was then
analysed as described above. The two procedures were carried out on the same urine sample and the difference between the two results gave the ethereal sulphate.

Creatinine concentration was determined by the method described previously (Ikeda and Ohtsuji, 1969a).

Results were in some instances adjusted to a specific gravity of urine of 1-016 as described by Elkins and Pagnotto (1965) or expressed in terms of creatinine concentration (Jackson, 1966).

**Phenol in air**

Two midget impinger tubes, each containing 15 ml of 0-1 M borate buffer (pH 10-0), were connected in series with a pump-battery-flow meter assembly ('portable pump', Mines Safety Appliances Co., Pittsburg, U.S.A.) and 5 to 10 l. of air were passed through them at about 2 l/min. A 3-5 ml portion of the buffer (diluted when necessary with fresh buffer) was taken from each tube, 0-1 ml of 0-2% 2,6-dichloroquinone chloroimide in ethanol (Gibbs' reagent) was added (Gibbs, 1927), and the resulting blue colour was measured at 610 nm 20 minutes after the addition. At least five determinations were carried out at various sites in each room, the average of which was taken to represent the environment.

**Statistical analyses**

Urinary data were assumed to be log-normally distributed (Heath, 1967) for statistical analyses. Regression lines were calculated with weighting for the numbers of samples (Snedecor, 1956).

**Results and discussion**

As observed in toluene metabolism (Ikeda and Ohtsuji, 1969b), the total urinary phenol concentration varied proportionately to the environmental phenol concentration (Fig. 1A, C, E). The regression lines cut the vertical axis above zero. It is well known that human subjects, even with no known exposure to phenol or its mother compounds, excrete phenolic compounds into urine (Williams, 1959), which may be derived from aromatic amino acids or food components (see also van Haaften and Sie (1965)).

The concentration of conjugated phenol (the main components of which are presumably phenyl glucuronide and phenyl sulphate, as observed by Parke and Williams (1953) in rabbits' urine) also

![Graph showing the relationship between phenol in air and in urine](https://example.com/graph.png)

**FIG. 1.** Relation between environmental phenol concentration in air and urinary phenols: (A) total and free phenols (observed values); (B) conjugated phenol (observed values); (C) total and free phenols (corrected to a specific gravity of 1-016); (D) conjugated phenol (corrected to a specific gravity of 1-016); (E) total and free phenols (corrected to constant creatinine concentration); (F) conjugated phenol (corrected to constant creatinine concentration).

Symbols show geometric means. Lines are weighted regression lines for the means (- - -) together with those for the S.D. ranges (-----).

The numbers of samples represented by each point were as follows: 0 mg/m³, 7; 0-6 mg/m³, 7; 4-2 mg/m³, 4; 7-8 mg/m³, 8; 8-8 mg/m³, 8; 9-6 mg/m³, 8; and 12-5 mg/m³, 7. Urine samples were collected at 3.00 p.m. towards the end of the day's exposure.
increased linearly with environmental phenol concentration. The concentration of free (i.e., unconjugated) phenol, on the contrary, was essentially independent of environmental phenol. This suggests that human subjects possess ample capacity to conjugate absorbed phenol as glucuronide or sulphate under the conditions studied, and perhaps up to the phenol concentration of 19 mg/m³, the A.C.G.I.H. threshold limit value for 1969.

From the results shown in Fig. 1C a tentative calculation was made to estimate the efficiency with which phenol is taken up through the lung. The regression line for total urinary phenol cuts the ordinate at 55 mg/l and reaches 262 mg/l when phenol in air is 10 mg/m³. The difference is attributable to phenol absorbed after inhalation. Because the half-life of phenol in man is short (Parke and Williams, 1953; also the data in Fig. 2), it may be reasonable to assume that the intake and excretion of phenol had come to a balance by the time urine was sampled at 3.00 p.m., as was the case in toluene metabolism (Ogata, Nagao, and Tomokuni, 1968). If we assume that the respiratory volume under working conditions was 15 to 20 l/min and that the urine output was around 50 ml/hr, the estimated amount of environmental phenol inhaled was about 10.5 mg/hr, i.e., 10 × (15 to 20) × 60/1,000, while the amount of urinary phenol attributable to phenol absorbed was 10.2 mg/hr, i.e., (262 – 55) × 50/1,000. This agreement between the two estimates, even though approximate, indicates that phenol was taken up very efficiently through the lungs.

Other examples are given in Fig. 2A and B to show that the urinary concentration of phenol reflects the exposure to environmental phenol. Urine samples collected at 8.00 a.m. (before exposure) on the sixth day of work after five consecutive working days contained only a little phenol. One working day with exposure caused a sharp increase in urinary phenol concentration, which returned to normal after the subjects were removed from the environment for two days (Fig. 2A) or even for one (Fig. 2B). Exposure to phenol on the next day again resulted in a rise of urinary phenol. These alterations were entirely attributable to the changes in concentration of conjugated phenol; consistent with the results given in Fig. 1, the levels of free phenol remained essentially unchanged regardless of exposure. The pre-exposure level on the sixth day of work in Fig. 2A was significantly (P > 0.05) higher than the level in the urine of controls when tested by t test, but this may not mean accumulation of phenol in the body during the working day because the corresponding pre-exposure level in Fig. 2B was almost in the normal range.

Neither urinary ethereal sulphate nor urinary ethereal glucuronide was a good index of phenol exposure. As shown in Figs. 3 and 4, the regression lines for both compounds have very small gradients and wide S.D. ranges. This poor validity of ethereal sulphate as an exposure index agrees with the poor reliability of the urinary sulphate ratio as an index of benzene exposure. The phenol test is now taking the

![Figure 2](http://oem.bmj.com/article-figs/1972-0101136-fig2.jpg)

**FIG. 2.** Changes in urinary concentrations of phenols after exposure to environmental phenol. Columns show geometric means. The shaded zone on the vertical axis shows fiducial limit (P = 0.95) ranges of normal values for total phenol (Ikeda and Ohtsui, 1969a).

(A) Four urine samples from men exposed to 8.8 mg/m³ phenol were collected at each time period, i.e., at 8.00 a.m. before exposure and at 3.00 p.m. Two days without exposure lay between the two days with exposure (shown with thick lines on the horizontal axis).

(B) Six urine samples from men at 7.8 or 9.6 mg/m³ phenol. One day without exposure lay between the two days with exposure.
Phenol concentrations in air and in urine

FIG. 3. Urinary excretion of ethereal sulphate from men exposed to phenol in air at various concentrations.

FIG. 4. Urinary excretion of ethereal glucuronide from men exposed to phenol in air at various concentrations.

The lines are weighted calculated regression lines (---) together with those for the S.D. ranges (---). The numbers of samples are as described in Fig. 1.

place of the sulphate ratio test as the environmental exposure to benzene lessens. (For a review, see Browning, 1965.) The highest concentration of phenol investigated here was 12-5 mg/m³, which is equivalent to 10.3 mg/m³ (3.3 p.p.m.) of benzene on an equimolar basis but less when the low efficiency with which benzene is absorbed (Nomiyama and Nomiyama, 1969) is borne in mind. According to Elkins (1954), benzene concentrations as high as 40 p.p.m. are necessary to change the sulphate ratio.

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References


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