Urinary thioether output as an index of occupational chemical exposure in petroleum retailers

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Petroleum is a complex mixture of lipophilic chemicals, some of which are likely to be substrates for the microsomal mixed function oxygenases. Petroleum retailers, by virtue of their occupational exposure potential, are a group likely to receive relatively heavy exposure to petroleum products in comparison with the general public.

Urinary thioether output may provide a simple, non-invasive technique for identifying occupational chemical exposure. The technique exploits the fact that conjugation with glutathione, followed by urinary elimination as mercapturates, is a significant metabolic clearance pathway for electrophilic chemicals and, in particular, for putatively toxic metabolites of the microsomal mixed function oxygenase system. Raised thioether output has been confirmed for several occupations associated with potential exposure to such chemicals.

Induction of the microsomal oxygenases is a common response to the ingestion of lipophilic chemicals. Since hepatic microsomes from rats exposed to petroleum vapours were found to have induced activity towards a range of cytochrome p450 substrates, it was reasonable to interpret the faster clearance of antipyrine in a cohort of petroleum retailers to result from occupational exposure to petroleum, although it was not possible to confirm exposure more directly using indices based on petroleum components such as benzene or lead and shown by others to be sensitive to petroleum exposure.

This study was undertaken to assess the utility of the urinary thioether technique to detect occupational chemical exposure in petroleum retailers. Since attendant operated pumps have been supplanted in many outlets by self-service pumps, it was of some interest to determine whether this difference in retailing practice was reflected in the occupational exposure patterns.

Methods

Forty three male and five female employees (aged 16–53 (mean 29, SD 10)) at 25 Adelaide suburban petrol vending stations took part in the study. Thirteen were employed in self service stations. All had worked in the industry as petrol pump attendants or garage mechanics, or both, for more than a year. They were asked to collect two urine samples during a midweek working day—one prework sample between 0600 and 0700 and one postwork sample between 1900 and 2000. Samples were stored in the freezer until analysis. A short questionnaire giving details of diet, medications taken, cigarette consumption, and employment duties was completed by each subject at the time of sample collection. Urine samples were analysed for both thioether and creatinine concentration as follows.

THIOETHER

Urinary thioether concentration was determined after acidification and ethylacetate extraction by colorimetric assay, as described by van Doorn et al. Concentrations were expressed as μmoles thioether relative to creatinine concentration (μmoles) to minimise any variation due to fluctuations in urine output.

CREATININE

Creatinine concentration was determined by the method of Yatzidis.

STATISTICS

Work related changes (am to pm differences) in individuals were analysed by the Wilcoxon matched pairs, signed rank test, 2 tail. The rises in urinary thioether output in smokers versus non-smokers, in pump attendants versus self service outlets, and the influence of workshop duties were compared with the Mann-Whitney U test. The interaction between smoking and work related thioether output was assessed by a 2 way ANOVAR.
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Results

Urinary thioether concentrations in petrol retailers at the end of a normal working day were significantly raised when compared with the morning sample (p < 0.001) (figs 1 and 2). The differences were greater (p < 0.001) in attendant operated outlets than in self service outlets (fig 1). Whereas most of the self service employees were solely engaged in remote operation of the cash register and pump controls, some also undertook mechanical workshop duties. In neither self service nor attendant operated stations, however, was there any indication that workshop duties were a significant factor in increasing urinary thioether output (fig 2). The only factor, other than driveway exposure to petroleum products, that could be identified as being associated with raised thioether output was cigarette smoking. Cigarette smokers excreted higher levels of thioethers in both prework (p < 0.005) and postwork (p < 0.001) samples, and there was also a significant interaction between smoking and work related thioether output (p = 0.01).

Discussion

These results confirm that raised urinary thioether output attributable to occupational chemical exposure occurs in the petroleum retailing industry, in common with findings in other industries.3–7 In the strict sense the technique should be viewed as a qualitative, rather than a quantitative, indicator of exposure. The estimated thioether concentrations are based on thiol determination after hydrolysis of an unknown mixture of thioethers in urine, and neither the identity of the metabolites, nor their quantitative recovery, can be assumed. While it has been shown that this non-specific technique detects dose related increases in output from certain model substrates in
animals,\textsuperscript{11–13} the human exposure model is more complex.

Given the non-specific nature of the assay, it is not possible to identify the chemical components of the occupational environment that account for the raised thioether output. Nevertheless, the observations that (1) raised thioether output is not associated with performance of workshop duties, and (2) isolation from driveway pump operation in self service outlets reduces the magnitude of the increase leads to the strong implication that inhalation of petrol vapour is the main source of thiomethylisable substrates.

Metabolic pathways leading to the formation of the urinary thioethers need not necessarily involve an electrophilic intermediate such as an epoxide of the microsomal oxygenases. Petroleum contains such substances as aliphatic and aromatic halides and alpha-beta unsaturated ketones, many of which undergo direct or glutathione transferase catalysed conjugation with glutathione. The direct and indirect pathways cannot be differentiated in our data, although the apparent interactive effect of cigarette smoking may favour an oxidative step in the dominant pathway on the grounds that smoking is known to induce microsomal oxygenase activity.\textsuperscript{14}

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References


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