Determinants of benzo(a)pyrenediol epoxide adducts to haemoglobin in workers exposed to polycyclic aromatic hydrocarbons

M Ferreira Junior, S Tas, M dell’Omo, G Goormans, J P Buchet, R Lauwerys

Abstract

Objective—The aim was to assess the determinants of benzo(a)pyrenediol epoxide adducts to haemoglobin (BaPDE-Hb) in workers exposed to polycyclic aromatic hydrocarbons (PAHs).

Methods—This was a study of the correlations between the concentrations of PAHs in air, 1-hydroxypyrene in urine, and BaPDE-Hb adduct concentration in the blood in 206 men working in steel foundries and one graphite electrode producing plant, taking into consideration confounding factors such as smoking and dietary habits.

Results—BaPDE-Hb adduct concentration was correlated (r = 0.26; p = 0.0002) with the airborne PAH concentration and was influenced by tobacco consumption but not by dietary habits. Benzo(a)-pyrene concentration in air, 1-hydroxypyrene concentration in urine, and duration of exposure to PAHs were not associated with the adduct concentration.

Conclusion—Although environmental exposure to PAHs was statistically associated with BaPDE-Hb adduct concentration, differences between individual subjects in the metabolism of benzo(a)pyrene probably play an important part in determining the amount of BaPDE-Hb adducts formed.

Key words: polycyclic aromatic hydrocarbons, benzo(a)pyrene, adducts to haemoglobin

Occupational exposure to mixtures of chemicals containing benzo(a)pyrene and several other polycyclic aromatic hydrocarbons (PAHs) has been associated with an increased mortality risk from neoplasms. Different approaches have been proposed for assessing occupational exposure to PAHs—namely, air analysis, skin contact monitoring (for example, by cutaneous pads), and biological analyses. Several studies have confirmed that 1-hydroxypyrene, a metabolite of pyrene excreted in urine, reflects recent PAH absorption by all routes.

As covalent binding of electrophilic PAH metabolites to DNA is considered as a critical step in the initiation of cancer, methods have been developed for the detection of PAH-DNA adducts in human tissues and peripheral white blood cells, with the aim of assessing the biologically effective dose and of estimating the risk at the individual level (see dell’Omo and Lauwerys for a review).

Because genotoxic compounds can react with haemoglobin and serum proteins to form stable adducts, assessment of PAH-blood protein adducts has been considered as an alternative marker of exposure to PAHs. Because the lifetime of haemoglobin in humans is about 120 days, and no enzymatic repair systems exist for haemoglobin, measurement of haemoglobin adducts might reflect the integrated exposure to PAHs over a period of four months before sampling. Therefore, interpretation of haemoglobin adduct measurement might be easier than that of DNA adducts. Despite these apparent advantages, there have been few reports on the detection of PAH-haemoglobin adducts in humans and they mainly focused on the analytical aspects. More data are therefore necessary to establish the real value of this biological marker for assessing human exposure to PAHs. We have measured benzo(a)pyrenediol epoxide haemoglobin (BaPDE-Hb) adducts in workers from two coke oven plants and one graphite electrode plant, and have attempted to identify the factors influencing their concentration.

Subjects and methods

STUDY POPULATION
A detailed description of the study population can be found in a previous publication. It involves 286 men working in steel foundries (mainly in two coke ovens and rolling mills) and one graphite electrode producing plant, but measurement of BaPDE-Hb adducts could only be performed on 206 subjects.

Answers from a self administered questionnaire inquiring about present health state, medical history, alimentary, smoking, and alcoholic habits, educational level, dwelling site, and occupational activities were obtained from each participant.
Each worker was equipped with a personal air sampling pump (model 224 PC3EXR3, SKC Inc, Eighty Four, PA, USA, flow rate of 2 l/min) during at least a five hour period of one normal workshift. Particles were retained on a glass microfibre filter (GFF, Whatman, 3-7 cm diameter) and vapours were adsorbed on a two section Chromosorb 102 tube placed between the pump and the filter. Tubes and filters were kept in a refrigerated dark room until analysis (within two weeks). At the end of the shift samples of blood and urine were collected for the determination of BaPDE-Hb adducts, 1-hydroxypyrene in urine, and other biological variables (see later).

Two criteria were used to classify the workers as being exposed to PAHs: (1) average PAH concentration in air > 4 μg/m³ or (2) 1-hydroxypyrene concentration in urine > 1 μg/g creatinine. These represent the 95th percentile values of the distributions of PAH concentration in air and 1-hydroxypyrene concentration in the urine of workers not occupationally exposed.7

Methods
Determination of PAHs in Air
Benzo(a)pyrene and 12 other PAHs (naphthalene, fluorene, phenanthrene, anthracene, fluoranthene, pyrene, benz(a)anthracene, chrysene, benzo(e)pyrene, perylene, dibenz-(a)anthracene, and benzo(g,h,i)perylene) were determined by high performance liquid chromatography (HPLC) coupled to fluorometry as previously reported.4 Total PAH was calculated as the sum of the 13 PAHs. The results were expressed as μg/m³ of air.

Determination of 1-Hydroxypyrene in Urine
The technique used was that of Jongeneelen et al.23 Results were standardised for urinary creatinine and expressed as μg/g creatinine.

Determination of BaPDE-Hb Adducts
The BaPDE-Hb adducts were determined by HPLC with fluorescence detection according to a procedure adapted from Shugart.24 Briefly it comprised (1) isolation of the red blood cells by centrifugation (2000 rpm for 10 minutes) and lysis of the cells by mixing with distilled water (v/v); (2) mild acid hydrolysis of the haemoglobin (0.1N HCL at 80°C for four hours) releasing benzo(a)pyrene (BaPtetrol) (only the isomer r7,t8,h10 BaPtetrol (tetrol 1–1) was detected); (3) clean up of the hydrolysate on successively, a C18 cartridge (Varian), diethylaminoethyl-cellulose (DE 32 Whatman), and Bond elut 3-ml phenyl (Varian) columns, elution with methanol, evaporation under a gentle stream of nitrogen, and redispersion in 500 μl of methanol; (4) HPLC/fluorescence detection. Results were calculated from a standard curve with tetrol 1–1 (Midwest Research Institute), standardized for the blood haemoglobin concentration, and expressed in fmol BaPtetrol/mg Hb. The recovery was assessed from r7,9,12-dihydroxy-9,10-epoxy-7,8,9,10-tetrahydrol[1,3 ³H] benzo(a)pyrene (Chemsyn Science Laboratories) as an internal standard.

Other Biological Analyses
Urinary creatinine was determined by the colorimetric method of Jaffé with slight modifications for automation (Technicon RA1000, Tarrytown, NY, USA).

Urinary thiocyanate concentration was measured by the technique of Pettigrew and Fell.25

Urinary ethanol was determined by gas chromatography on a 150 cm, 3-2 mm internal diameter column containing 25% Carbowax 20M on Chromosorb WAW DMCS 100–120 mesh at 90°C coupled with a flame ionisation detector.

The hepatic enzymes glutamic oxaloacetic transaminase (S-GOT), serum glutamic pyruvic transaminase (S-GPT), and γ-glutamyl transpeptidase (γ-GT) were measured with Biochemica test combination kits (Boehringer Diagnostica, Mannheim GMBH, Germany).

Haemoglobin was analysed by a cyanohaemoglobin procedure.

Serum creatinine was determined by a direct colorimetric method.

Urinary albumin, retinol binding protein (RBP), and β₂-microglobulin (β₂M) were determined by automated non-isotopic immunonasays based on latex particle agglutination.26

Urinary N-acetyl-β-D-glucosaminidase (NAG) was measured by the fluorimetric method described by Tucker et al.27

Statistical Analysis
Responses to the questionnaire and analytical data were stored in a database and statistically analysed with SAS/STAT software.28 When necessary (Wilk’s test), distributions were normalised by log transformation.

The statistical significance for differences between means was assessed by Student’s t test.

A stepwise regression analysis was performed to identify the statistically significant determinants of BaPDE-Hb adducts. As well as the exposure variables (benzo(a)pyrene or total PAHs in air or 1-hydroxypyrene in urine) the various determinants introduced in the model were age, urinary thiocyanate, urinary ethanol, number of cups of coffee daily, risk food index (0 = no regular consumption of grilled and fried foods; 1 = regular consumption of these food items), protective food index (0 = no regular consumption of fruits and juices; 1 = regular consumption of these food items), dwelling site (0 = urban area; 1 = countryside), plant (0 = coke oven; 1 = graphite electrode), renal and hepatic indices (0 = none of the renal variables (serum creatinine, urinary albumin, RBP, NAG, β₂M) or the serum hepatic enzymes (S-GOT, S-GPT, and γ-GT) exceeding the 95th percentile of the corresponding distribution in the control group; 1 = at least one renal or hepatic variable above this value). The procedure ended...
Table 1  Concentration of BaPDE-Hb adducts (fmol/mg Hb) according to the concentration of PAHs in air and 1-hydroxypyrene (1-HOP) in urine

<table>
<thead>
<tr>
<th>Exposure variable</th>
<th>No of subjects</th>
<th>BaPDE-Hb adducts</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PAHs in air (µg/m³):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤4-0</td>
<td>113</td>
<td>24-3 (2-5)*</td>
<td>0-03</td>
</tr>
<tr>
<td>&gt;4-0</td>
<td>93</td>
<td>31-8 (2-3)</td>
<td>0-01</td>
</tr>
<tr>
<td>1-HOP in urine (µg/g creatinine):</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>≤2-0</td>
<td>139</td>
<td>27-2 (2-4)*</td>
<td>NS</td>
</tr>
<tr>
<td>&gt;2-0</td>
<td>64</td>
<td>27-7 (2-5)</td>
<td>0-01</td>
</tr>
</tbody>
</table>

*Geometric mean (SD). †Range. ‡Three workers failed to give a urine sample at the end of the workshift.

Table 2  Determinants of BaPDE-Hb adducts (multiple regression analysis) in the total population

<table>
<thead>
<tr>
<th>Variable</th>
<th>Determinants *</th>
<th>Partial regression coefficient</th>
<th>Partial correlation coefficient (r*)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>BaPDE-Hb adducts</td>
<td>PAHs in air</td>
<td>-0-107</td>
<td>0-062</td>
<td>0-003</td>
</tr>
<tr>
<td></td>
<td>Thiocyanate in urine</td>
<td>-0-304</td>
<td>0-031</td>
<td>0-01</td>
</tr>
<tr>
<td></td>
<td>Renal index</td>
<td>-0-018</td>
<td>0-024</td>
<td>0-02</td>
</tr>
</tbody>
</table>

* Determinants included in the regression models are: benzo(a)pyrene or total PAH concentration in air or 1 hydroxypyrene in postshift urine, thiocyanate in urine, ethanol in urine, coffee consumption (number of cups/day), risk food index (fried and grilled foods), protective food index (fruits and vegetables), age, dwelling site, renal index, hepatic index, and type of plant.

Results

The mean concentration of BaPDE-Hb adducts was significantly increased in workers (n = 93) with a time weighted average exposure to PAHs above 4-0 µg/m³ (table 1). No significant difference between adduct mean concentrations was found when the 1-hydroxypyrene concentration in urine was used as the exposure criterion. The multiple regression analysis performed on the total population identified concentration of PAHs, thiocyanate concentration in urine (used as biological indicator of smoking habits), and renal index as statistically significant determinants of BaPDE-Hb adduct concentration (figure; table 2). The 1-hydroxypyrene concentration in urine was not associated with the concentration of BaPDE-Hb adducts and surprisingly the time weighted average exposure to BaP alone was not a significant determinant.

The possible effect of the duration of occupational exposure to PAHs (mean = 10-5 (range 0-25-30-9) years) on the concentration of BaPDE-Hb adducts was tested with linear regression models including the variables mentioned. The models were applied to the workers exposed to PAHs, classified either on the basis of PAH concentration in air (n = 93) or 1-hydroxypyrene concentration in urine (n = 64). Neither model showed duration of exposure as a significant determinant of BaPDE-Hb adduct concentration.

In the logistic regression model, none of the exposure variables (PAHs in air, benzo(a)pyrene in air, 1-hydroxypyrene in urine, duration of exposure) emerged as a significant determinant of an increased prevalence of increased values (> 70 fmol/mg Hb) of BaPDE-Hb adducts (results not shown).
Discussion
So far, no large scale epidemiological study has been carried out involving the monitoring of PAH-Hb adducts in workers exposed to PAHs. We have previously reported that in the plants in which the present study was carried out, the proportion of the measured airborne PAHs (particularly the sum of the three carcinogenic compounds, benzo(a)anthracene, benzo(a)pyrene, and dibenz(a,h) anthracene) was similar and correlated with 1-hydroxypyrene in urine. Therefore, benzo(a)pyrene was selected as the prototype of PAHs and we have attempted to determine whether BaPDE-Hb concentration was associated with some exposure variables (benzo(a)pyrene and total PAHs in air, 1-hydroxypyrene in urine, duration of exposure) and possible confounders (smoking and dietary habits, residence, renal, and hepatic function).

We noted that when the total PAHs in air was used as the criterion of exposure, mean BaPDE-Hb adduct concentration was higher in the exposed than in the control workers. This is in line with the higher concentration of PAH-protein adducts found by Shersen et al.20 in exposed foundry workers. The multivariable regression analysis however, showed that although the total concentrations of PAHs in air was the main determinant of BaPDE-Hb adduct concentrations, it only explained 6% of their variance. Smoking habits (assessed by the thiocyanate concentration in urine)29 also influenced the amount of adduct detected, but no significant association was found with 1-hydroxypyrene in urine (an indicator of the overall amount of PAHs recently absorbed), benzo(a)pyrene in air, or duration of exposure. It should be noted that some authors21 also did not find an association between PAH-DNA adducts in peripheral lymphocytes and total PAHs in air or 1-hydroxypyrene in urine. One has to keep in mind that both benzo(a)pyrene in air and 1-hydroxypyrene in urine reflect recent exposure whereas BaPDE-Hb adducts may integrate exposure over a four month period. The impact of tobacco consumption on BaPDE-Hb adduct concentrations was also reported by Wester et al.21 resulting from the presence of PAHs in tobacco smoke, increased occupational exposure to PAHs due to smoking at the workplace, or possible interference of tobacco smoke on benzo(a)pyrene biotransformation by enzymatic induction. As renal function declines with age, the negative association between the renal index and the BaPDE-Hb adduct concentration could be due to a selection bias; the more polluted workplaces in the plants might have been allotted to the youngest workers. This explanation is unlikely as age was not indentified as a confounder in the multiple regression model tested and no significant difference was found between the mean age of exposed workers with or without abnormal renal variables. It should be noted, however, that none of the renal markers was specifically responsible for this association.

In conclusion, the results suggest that although environmental exposure to PAHs is statistically correlated with the amount of BaPDE-Hb adducts (figure), it is not the main determinant as it only explains 6% of their variability. It is possible that endogenous toxicokinetic factors (interindividual differences in the metabolism of benzo(a)pyrene)13 may be critical in determining the amount of haemoglobin adduct formed. The interest in BaPDE-Hb adduct measurement to reflect the biologically active dose of benzo(a)pyrene remains to be established. Assessment of their relation with markers of genotoxic effects (for example, sister chromatid exchanges and chromosomal aberrations in peripheral lymphocytes), and of susceptibility may help clarify this question.

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