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Bladder cancer mortality of workers exposed to aromatic amines: an updated analysis


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Bladder cancer mortality of a cohort of dyestuff workers at a factory in the province of Turin, Northern Italy, was followed up to 1981, when 41 deaths from bladder cancer were observed. Less than one had been expected (Decarli et al., 1985). One of the open questions in that analysis concerned the pattern of relative and absolute excess risk after stopping exposure, since the use of known carcinogenic amines in dyestuff production was stopped in 1972 (Rubino et al., 1982). This would have important implications as regards our understanding of the process of carcinogenesis by aromatic amines, particularly their potential impact on one of the latter stages of the process (promotion) (Day & Brown, 1980). A better knowledge of the risk following cessation of exposure would also have clear public health relevance, since it could provide indications for optimising follow-up schemes for ex-workers.

In this analysis, we have thus added 8 years of follow-up to the same cohort. Briefly, the dataset comprised all men who had been employed since 1946, and had worked for at least 1 year in the factory between 1922 and 1970. For the 906 workers meeting these criteria, date of birth, of employment(s) and termination of employment(s), the last known address, and detailed job particulars including categories of exposure to selected chemicals were abstracted from the factory’s personnel records. Only 38 subjects were lost to follow-up. Further, 204 workers not directly involved in exposure to aromatic amines were excluded. Death certificates were obtained from registration offices in the municipality of death, and further verification of vital status was obtained from registries of current residence. Causes of death were coded according to the International Classification of Diseases (ICD). No further detail was available on tumour characteristics, or on potentially relevant covariates, such as smoking. Further details on exposure classification and follow-up are given in Decarli et al. (1985). For the present report, follow-up was updated to December 1989, for a total number of 271 deaths among 664 exposed subjects (49 from bladder cancer) and 19,157 man-years at risk (14,570 among exposed subjects).

The analyses presented are centered on patterns of risk with time since stopping exposure among the 664 subjects exposed to aromatic amines. As in the previous report, they are based on simple comparisons of observed and expected (based on national mortality rates (La Vecchia et al., 1990)) numbers of deaths, and on fitting two general models of risk (multiplicative, or relative risk; additive, or excess risk (Baker & Nelder, 1978; Breslow & Day, 1987)), to allow simultaneously for various related factors (age at first exposure; duration of exposure; job category; time since last exposure).

Table I gives numbers of observed and expected deaths from bladder cancer, and all other causes of death, according to time since last exposure. Overall, there were 49 bladder cancer deaths vs 1.6 expected, corresponding to a standardised mortality ratio (SMR) of 30.4. A total of 222 deaths from other causes were observed vs 161.8 expected (SMR 1.4). Rates were elevated for upper digestive and respiratory tract neoplasms (oral cavity, six observed vs 2.2 expected; oesophagus, four observed vs 1.7 expected; larynx, nine observed vs 2.4 expected), and other alcohol-related causes. In the descriptive analysis, a clear trend of decreasing risk with longer time since last exposure was observed, both for bladder cancer (from 100.8 during exposure to 14.8 20 years or more after the last exposure) and for all other causes (from 1.9 to 0.9).

Table II gives the parameter estimates from the multiplicative (relative risk) and additive (absolute excess risk) models, under which the effects of the three time factors, age at first exposure, duration and time since last exposure (plus job category, expressed as type and degree of exposure to selected carcinogens, Rubino et al., 1982) were estimated simultaneously. These estimates, together with the corresponding standard errors, are expressed in relation to one of the categories of each variable, arbitrarily chosen as referent; the exponential of each estimate gives the risk for the corresponding category. Statistical significance can be tested by comparing the ratio of each parameter estimate to its standard error with a standardised normal deviate.

The relative risk was strongly inversely related to age at first exposure, whereas the absolute excess risk was unrelated to it. Duration of exposure showed a strong direct association with absolute excess risk but no significant relationship with relative risk. Time since last exposure was inversely related to relative risk, but not to absolute excess risk, which showed no significant value up to 20 years after stopping exposure. In relation to job category, manufacture of a-β naphthylamine or benzidine was associated to the highest risk, followed by fuchsian or safranine T manufacture and by use or intermittent exposure to naphthylamine or benzidine. The association with job category was of similar magnitude using the multiplicative and the additive model.

Some of these results are interpretable, as previously discussed (Decarli et al., 1985), in terms of the multistage theory of carcinogenesis (Armitage & Doll, 1961; Day & Brown, 1980). For instance, the inverse relation of the relative risk with age at first exposure and the absence of association with absolute excess risk are compatible with an early-stage effect of aromatic amines on bladder carcinogenesis. The apparent anomaly of the absence of association of relative risk with duration may be due to the fact that relative risk is a function of (d/t), where d is duration and t is age (and hence the sum of age at first exposure, duration and time since last exposure) (Brown & Chu, 1983; Day & Brown, 1980).

The contrasting results for relative and absolute excess risk
with reference to time since last exposure are more important, particularly since this is the variable for which most information has been added in this updated analysis. The inverse relationship of the relative risk with time since last exposure is of interest from an etiological viewpoint and indicates the existence of a late stage effect in the process of aromatic amine carcinogenesis, besides the early stage one suggested by the pattern of relative and absolute excess risk with age at first exposure.

However, although the multiplicative model shows a decreasing function of the relative risk with longer time since last exposure, the additive model indicates that the absolute excess risk flattens off but does not decline after stopping exposure – even 20 years or more since last exposure – and, in fact, eight additional deaths were observed during 8 further years of follow-up. This shows interesting similarities with the pattern of risk observed after stopping smoking, with a levelling of lung cancer mortality around the levels reached at the time of stopping, which was interpreted as indicative for the existence of a late (penultimate) stage effect of tobacco on bronchial carcinogenesis (Doll, 1971; Armitage, 1971). Thus, the present data are consistent with the time risk relationship for cigarette smoking and bronchial carcinoma, where lung cancer incidence is a simple power function of total duration of smoking independently from age at starting or any other time related factor (Doll & Peto, 1978).

Further, the pattern of absolute excess risk after stopping has major public health relevance, since it stresses the importance of long-term continued surveillance for bladder cancer in this cohort.

This underlines the complementarity of the information conveyed by the two models, and the interest of their implications on a theoretical (carcinogenesis) and practical (public health) level (Breslow & Day, 1987), besides the importance of a long-time follow-up for defining the ultimate impact of aromatic amines on bladder cancer risk.

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References


