Correspondence

Pulmonary fibrosis in asbestos insulation workers with lung cancer

SIR,—There has been discussion of the paper of Kipen et al (1987;44:96-100) by Rudd1 and a response by Suzuki et al2 but neither of these letters has dealt with the basic flaw in the design of the original investigation.

Kipen et al reviewed 138 cases of histologically confirmed lung cancer among 450 diagnosed in the United States-Canadian insulator cohort. The restriction to 138 was because only this number had adequate chest x-ray films and adequate samples of pulmonary tissue for microscopic evaluation. Whether the 138 cases were representative of all the cases is open to question but let us assume that they were. The major problems were: (1) it was assumed that all interstitial fibrosis, even with rare or no asbestos bodies, was asbestosis by the authors since they equated the two terms in two places; (2) there were no unexposed control groups with “blind” evaluation of either the chest x-ray films or the histological specimens.

Regarding the first problem, it must be recognised that agents other than asbestos cause interstitial fibrosis, perhaps most commonly cigarette smoke, and asbestosis has no pathognomonic x-ray or histological signs. Therefore, the second problem, the lack of “blindly assessed” controls, becomes the most serious defect in this study. Without “blindly evaluated” controls, it is impossible to assess the contribution of asbestos to the causation of lung cancer in asbestos workers.

Among the United States-Canadian insulators investigated, the relative risk of lung cancer 20 or more years after onset of exposure may be calculated to be 4.23 from data in the paper by Selikoff et al4 using death certificate diagnoses to provide a valid comparison with the risk in the general population. From this relative risk, it is obvious that 23.6%—that is, one out of every 4.23 cases of lung cancer—would have occurred in this population without exposure to asbestos. I made this calculation based on the interval of 20 or more years from onset of exposure because 134 of the 138 cases studied by Kipen et al occurred 20 or more years from onset of exposure (data from their fig 3). This figure of 23.6% approximates the 18% figure in the series of 138 cases reported as having no radiographic signs of asbestosis in the pulmonary parenchyma. In view of the potential for error in the methods used, this approximation is remarkably good.

Since both asbestos and asbestos-related lung cancer have a dose response relation to asbestos, the most parsimonious interpretation of these juxtaposed data is that the cases of lung cancer without x-ray evidence of asbestosis are the cases that would have occurred in this cohort if there had been no exposure to asbestos—that is, they were the expected number. The conclusion by Kipen et al that their findings “indicate the primacy of the history of exposure to asbestos, irrespective of the presence or absence of non-malignant x-ray changes (asbestosis) when considering lung cancer possibly associated with occupational exposure to asbestos” is not warranted by the design of their study.

There is considerable circumstantial evidence that the risk of lung cancer is raised only among workers exposed to asbestos who also have parenchymal asbestosis. If this issue is ever to be settled definitively adequately designed studies must be undertaken.

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References


Incidence of cancer of the scrotum, 1971–84

SIR,—In 1984 Waldron et al reported on scrotal cancer in the West Midlands for the period 1936–76 (1984;41:445-9). They noted that the annual number of registrations appeared to be declining in the 1970s, compared with the relatively high numbers recorded in the 1960s. We can confirm that this trend has persisted having examined recent cancer registration data for the West Midlands Region (see figure). A recorded occupation was available for 109 (73%) of the 149 cases registered in the period 1970–84. Of these 109 occupations, at least 70% probably involved exposure either to mineral oils, pitch, or tar.

Annual numbers of registrations for England and
Wales are also shown in the figure. The downward trend is not due to changes in the size of the population or its age structure, since standardised registration ratios—calculated on the basis of the 1971 age specific rates—followed an almost identical pattern. A considerable proportion of cases of scrotal cancer are probably caused by exposures to chemicals, and the decline in incidence is, therefore, a likely consequence of improvements in occupational hygiene introduced in engineering industries over the past 20 years. It would be unreasonable, though, to assume that no new cases arise from current working practices.

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References


Influence of design characteristics on the outcome of retrospective cohort studies

SIR,—I read with interest this article by Swaen and Meijers (1988;45:624–9). They report that 76% of the retrospective cohort studies conducted by governmental agency epidemiologists are positive compared with 36% of the studies conducted by industry based epidemiologists. When the source of funding was considered, 77% of the studies funded by governmental agencies or universities were positive compared with 53% of the studies funded by industry. Swaen and Meijers indicate that studies conducted in the chemical industry are more likely to be negative than those conducted in other industries, irrespective of the funding source.

In discussing their research Swaen and Meijers state that "one possible explanation for this finding may be that those industries that employ epidemiologists are also more aware of the potential risks involved and have put greater effort into improving the occupational environment." I agree that this represents a possible explanation for the proportionally lower number of positive retrospective cohort studies of workers employed in the chemical industry or conducted by industry based epidemiologists. Those companies with inhouse epidemiology also tend to be those with long standing occupational medicine, industrial hygiene, and toxicology programmes. The more favourable health statistics suggest that the programmes have been effective in protecting employees.

I believe, however, that Swaen and Meijers failed to consider other alternative explanations for their findings. For example, they fail to consider the underlying reasons for undertaking a study. Academic and government epidemiologists may be more likely to conduct ad hoc studies. Often these studies are triggered by a pre-existing concern about an excess of disease in an occupational group. Such studies can arise because of a cluster of cases of a rare disease, unusual findings of a medical surveillance programme, or the results of other positive studies of the same or similar occupational exposures. These studies often quantify an association between an exposure and a health effect already supported by other data.

By contrast, several retrospective cohort studies conducted by industry based epidemiologists (or funded by industry) are mortality surveillance studies. These mortality surveillance studies are a part of company supported occupational health programmes and are conducted to help assure that safeguards implemented to protect the health of employees are adequate. In that context, it is not surprising that research findings are negative.

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