Radon and risk of extrapulmonary cancers: results of the German uranium miners’ cohort study, 1960–2003

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Data from the German miners’ cohort study were analysed to investigate whether radon in ambient air causes cancers other than lung cancer. The cohort includes 58 987 men who were employed for at least 6 months from 1946 to 1989 at the former Wismut uranium mining company in Eastern Germany. A total of 20 684 deaths were observed in the follow-up period from 1960 to 2003. The death rates for 24 individual cancer sites were compared with the age and calendar year-specific national death rates. Internal Poisson regression was used to estimate the excess relative risk (ERR) per unit of cumulative exposure to radon in working level months (WLM). The number of deaths observed (O) for extrapulmonary cancers combined was close to that expected (E) from national rates (n = 3340, O/E = 1.02; 95% confidence interval (CI): 0.98–1.05). Statistically significant increases in mortality were recorded for cancers of the stomach (O/E = 1.15; 95% CI: 1.06–1.25) and liver (O/E = 1.26; 95% CI: 1.07–1.48), whereas significant decreases were found for cancers of the tongue, mouth, salivary gland and pharynx combined (O/E = 0.80; 95% CI: 0.65–0.97) and those of the bladder: (O/E = 0.82; 95% CI: 0.70–0.95). A statistically significant relationship with cumulative radon exposure was observed for all extrapulmonary cancers (ERRWLM = 0.014%; 95% CI: 0.006–0.023%). Most sites showed positive exposure–response relationships, but these were insignificant or became insignificant after adjustment for potential confounders such as arsenic or dust exposure. The present data provide some evidence of increased risk of extrapulmonary cancers associated with radon, but chance and confounding cannot be ruled out.

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Although it is well established that occupational exposure to the radioactive gas, radon (222Rn), and its progeny increases the risk of lung cancer (BEIR, 1999; Laurier et al, 2004; Tomasek, 2004; Brüske-Hohlfeld et al, 2006; Grosche et al, 2006; Villeneuve et al, 2007; Tomasek et al, 2008; Vacquier et al, 2008), little is known about any effects on other cancers (Tomasek et al, 1993; Darby et al, 1995a; BEIR, 1999; Moehner et al, 2006; Rericha et al, 2006). As it is estimated that doses from radon and its progeny to organs other than the lung are approximately ≥100 times lower (Kendall and Smith, 2002; Marsh et al, 2008), large-scale occupational radon studies are required to investigate this possible relationship.

The largest and most informative study on this subject to date is the pooled analysis of 11 miners’ cohorts published by Darby et al (1995a). Overall, no statistically significant exposure–response relationship was observed, except for pancreatic cancer. The researchers concluded that high concentrations of radon in the air do not cause a material risk of mortality from cancers other than lung cancer. However, low statistical power, missing information on potential confounders and heterogeneity among the 11 studies were of concern. The aim of these analyses of the German Wismut uranium miners’ cohort study is to further evaluate the relationship between radon and extrapulmonary cancers. Compared with the pooled study, the Wismut cohort has a comparable size (58 987 vs 64 209), a longer average follow-up period (35 vs 17 years), a larger number of deaths from cancers other than lung cancer (3340 vs 1253), a longer mean duration of employment (12 vs 6 years) as well as a higher average cumulative exposure to radon (279 vs 155 working level months (WLM)). Moreover, information on occupational exposure to external γ-radiation, long-lived radioactive nuclides (LRNs), arsenic, fine dust and silica dust is available.

MATERIALS AND METHODS

Cohort definition and follow-up

The cohort has been described earlier (Grosche et al, 2006; Kreuzer et al, 2002, 2006). In brief, it represents a stratified random sample of 58 987 males, employed for at least 6 months from 1946 to 1989 at the former Wismut uranium company in East Germany. The first follow-up ran up to the end of 1998 (Grosche et al, 2006; Kreuzer et al, 2006), and this study extends the follow-up by 5 years through 2003. Information on the vital status of individuals was obtained from local registration offices, whereas death certificates were obtained not only from the responsible Public Health Administrations but also from central archives and the pathology archive of the Wismut company. The underlying causes of death from death certificates or the autopsy files were coded
follow-up (31 December 2003). The expected mortality rate was calculated by applying national mortality rates, grouped by calendar year and 5-year age bins, to the person-years in the grouped cohort data. The standardised mortality ratio (SMR) is given by the ratio of O/E, where O is the number of observed deaths in the cohort and E is the number expected from external rates. In common with other miners’ studies (Tomasek et al., 1993; Darby et al., 1995a), a 5-year lag was used to calculate the cumulative exposure to radon for all sites of cancer other than leukaemia and a zero lag for leukaemia. The confidence intervals of the SMR were calculated on the basis of the poisson distribution (Breslow and Day, 1987). SMRs were corrected for missing causes of death by dividing O by the proportion of known causes of death, P, which was binomially distributed. Different correction factors were used to account for the variability of P as, when methods were applied to account for this variability (Rittgen and Becker, 2000), the resulting SMR confidence intervals were not significantly affected.

The cancer sites examined were defined according to the pooled study by Darby et al. (1995a), but the 10th ICD code was used instead of the 9th ICD code. Earlier revisions of the ICD (8 and 9) and the former codes of the German Democratic Republic were recoded to ICD-10. As no separate external rates had been available for the time period 1968–1979 for cancers of the tongue and mouth, salivary gland and pharynx, these cancers were combined in the external analyses. In a few cases, the external rates were not available for certain years and cancer types and hence were not included in the corresponding external analyses, which is why the total numbers are sometimes lower than those in the internal regression and do not sum to the total number of non-lung cancers. A separate analysis for time periods < or > 10 years since first employment was performed, because earlier studies (Tomasek et al., 1993; Darby et al., 1995a) showed a lower mortality during the first period compared with the later period, most probably because of the selection of healthy men for employment in the mines. Owing to the long follow-up period in this cohort and the restriction of the follow-up period to 1960 and later, the proportion of cases occurring <10 years after the first employment was extremely low (1.7%) and thus did not affect the overall risk estimate.

Poisson regression was used to test for an association between cancer mortality risk and cumulative radon exposure. Tabulations of person-years at risk and cancer deaths were created with the DATABASE module of the EPICURE software (Preston et al., 1998). Cross-classifications were made by age, a, in 16 categories (<15, 15–19, 20–24, …, 85 + years), individual calendar year, y, in 58 categories and cumulative radon exposure, w, in seven categories (0, >0–49, 50–99, 100–499, 500–999, 1,000–1,499, 1,500 + WLM). The WLM categories were defined to be comparable with other studies (Darby et al., 1995a; Moehner et al., 2006), but with an added category of 0 WLM. The tabulated data were fitted to the following model – if r(a, y, w) is the age, year and exposure-specific cancer mortality rate and r(a,y) = r(a,y,0) is the baseline disease rate for non-exposed individuals, w = 0 then

\[ r(a,y,w) = r_0(a,y) \times \{1 + ERR(w)\} \]  

where ERR is the excess relative risk. A linear form for \( ERR(w) = bw \), with no dependence of the slope \( b \) on \( a \) and \( y \), was used to investigate the exposure–response relationship. In addition, a categorical analysis of the form \( ERR(w) = \sum_j \beta_j w_j \) was performed, where \( j \) refers to the exposure class. To test for the five potential confounders, LRN, internal \( \gamma \)-radiation, fine dust, quartz fine dust or arsenic, each of these variables \( z_i \), \( i = 1,5 \) was added separately to the model (1) with \( ERR(w, z_i) = bw + z_i \). Maximum likelihood with the AMFIT module of the EPICURE software (Preston et al., 1998) was used for estimation of the fit parameters: \( b, \gamma_i, \beta_j \) (\( j = 1,7 \)), and the internal baseline rates in
strata. Internal regression analyses were restricted to individual cancer sites with a total of >35 deaths.

RESULTS

In the follow-up period 1960–2003, a total of 57 199 persons were under observation, resulting in 1 762 208 person-years at risk and a mean duration of follow-up of 35 years. By the end of 2003, 35 294 (61.7%) men were alive, 20 684 (36.2%) had died, 233 (0.4%) had emigrated and 988 (1.7%) were lost to follow-up. The underlying cause of death was available for 19 501 (94.3%) of the deceased individuals. Some time during Wismut employment, whereas 7931 had never been exposed (Table 1). Those exposed received a mean cumulative exposure to radon of 279.4 WLM (median = 17 554) in dust-years (fine dust, silica dust). Figure 1 shows the annual mean exposure values for radon and its progeny in working level months (WLM), and for external γ-radiation in mSv and LRN in kBq m−3 for the exposed cohort members. Radon concentrations decreased sharply after 1955 because of the introduction of several ventilation measures, which led to conditions in accordance with the international radiation protection standards after 1970. In contrast to this, external γ-radiation and LRN show a different pattern, because their concentration was not affected by the improved ventilation. The annual mean exposure values for fine dust, silica dust and arsenic are given in Figure 2. Owing to the use of dry drilling, the concentrations of radon had been very high until 1955 and then decreased steadily with the implementation of wet drilling, reaching very low levels after 1970. A total of 17 554 miners were exposed to arsenic, with higher annual values in the early years compared with the later years.

Table 2 gives the numbers of O and E deaths based on the male Eastern German population, as well as the corresponding SMRs (O/E) with 95% CIs for all cancers other than lung cancer combined and for 24 individual cancer sites. The number of non-lung cancer deaths combined was close to expectation (O/E = 1.02; 95% CI: 0.98 – 1.05). Among 24 individual non-lung cancer sites, a significant excess was found for stomach (O/E = 1.15; 95% CI: 1.06 – 1.25) and liver cancers (O/E = 1.26; 95% CI: 1.07 – 1.48), as well as a significant deficit of cancers of the tongue, mouth, pharynx and salivary gland combined (O/E = 0.80; 95% CI: 0.65 – 0.97) and those of the bladder (O/E = 0.82; 95% CI: 0.70 – 0.95). Overall mortality was significantly higher than in the general population (O/E = 1.03; 95% CI: 1.02 – 1.05), mainly because of lung cancer (O/E = 2.03; 95% CI: 1.96 – 2.10).

In the internal regression analyses shown in Table 3, there is a significantly increased mortality from all cancers other than lung cancer with cumulative radon exposure (ERR/WLM = 0.014%; 95% CI: 0.006 – 0.023%). The two highest exposure categories 1000–1499 WLM and >1500 WLM show a 1.2-fold (95% CI: 1.02 – 1.38) and 1.16-fold (95% CI: 0.94 – 1.76) higher risk compared with the reference category of 0 WLM, respectively. Among the 18 individual sites with >35 cases, a significant positive relation with radon is observed for stomach cancer (ERR/WLM = 0.021%; 95% CI: 0.0007 – 0.043%), whereas excesses with borderline statistical significance were found for cancers of the pharynx (ERR/WLM = 0.16%; 95% CI: 0.045 to 0.37%) and liver (ERR/WLM = 0.044%; 95% CI: 0.008 to 0.096%). No association between leukaemia and cumulative radon exposure is found.
This is also true for all leukaemia except chronic lymphatic leukaemia (non-CLL) (n = 87, ERR/WLM = 0.019%; 95% CI: −0.04 to 0.088%), CLL (n = 40, ERR/WLM = −0.013%; 95% CI: −0.067 to 0.040%) and acute myeloid leukaemia (n = 31, ERR/WLM = 0.036%; 95% CI: −0.076 to 0.149%).

Overall, there is a low correlation between exposure to radon and exposure to external γ-radiation, LRN or arsenic (R < 0.28), whereas fine dust (R = 0.57) and silica dust (R = 0.63) are relatively highly correlated with radon exposure. Additional adjustment for each of the five factors showed no substantial modifying effect on the overall ERR/WLM for all non-lung cancers combined. In contrast, the adjustment led to a decreased risk for certain sites (e.g., stomach, larynx and liver) (Table 4). Overall, none of the risk estimates for the different cancer sites were significantly different after adjustment for the potential confounders.

**DISCUSSION**

In this study, a statistically significant relation between cumulative radon exposure and risk of extrapulmonary cancers combined is observed (ERR/WLM = −0.014%). After adjustment for potential confounders, such as exposure to arsenic, dust, LRN and γ-radiation, the ERR/WLM is only marginally modified, values of the ERR/WLM ranging from 0.016 to 0.011%, with some of the borderline significance. No earlier miners’ studies have reported a statistically significant result for this relationship (Tomasek et al., 1993; Darby et al., 1995a; Vacquier et al., 2008), and hence a non-causal chance result in our study cannot be ruled out. However, the earlier studies may have been limited by low statistical power. For example, in the pooled study by Darby et al. (1995a) an ERR/WLM of 0.01% for the time period >10 years after employment was observed, in line with our findings, but it is not statistically significant (Table 5). In both studies, an excess of non-lung cancers seems to be present only for exposure categories above 1000 WLM.

Dosimetric calculations indicate that extrapulmonary organs received very low doses compared with those received by the lung (Kendall and Smith (2002); Marsh et al., 2008). Marsh et al. (2008) recently estimated the absorbed doses for specific organs for several exposure scenarios in mines. For example, wet drilling, medium ventilation and medium physical activities were associated with the following doses in mGy/WLM: bronchial region 7.3, red bone marrow 0.031, kidney 0.02 and liver 0.0065. In our analyses, the ERR/WLM for lung cancer is approximately 14 times higher (n = 2999; ERR/WLM = 0.20%; 95% CI: 0.16–0.22%) than for non-lung cancers (n = 3,340; ERR/WLM = 0.014%), which is compatible with the biokinetic models. For individual sites, the majority showed a positive exposure–response relationship (15 from 18), although this was significant only for stomach cancer (Figure 3). After adjusting for the five potential confounders, however, no individual sites showed a significant exposure–response relationship.

**Specific sites**

**Liver** The increased mortality of liver cancer in miners compared with the general population (n = 158, O/E = 1.26; 95% CI: 1.06–1.25) is consistent with other miners' studies (Tomasek et al., 1993; Darby et al., 1995a,b) and appears not to be a chance finding. It may be because of the high consumption of alcohol among miners, which, in the early years, was offered (with cigarettes) free of charge. Alcohol abuse, or cirrhosis, was mentioned on the death certificate for 8%, or 37%, of the liver cancers, respectively. The principal two studies provided no evidence of a relationship with radon concentrations, however, no individual sites showed a significant exposure–response relationship.
Table 3  Relative risk for selected cancer sites by cumulative radon exposure based on internal poisson regression, 1960–2003

| Cancer site (ICD-10 code) | Cumulative radon exposure in working level months (WLM) | ERR/100 WLM | P-value |
|---------------------------|------------------------------------------------------|-------------|---------|
|                           | 0          | 0–49 | 50–99 | 100–499 | 500–999 | ≥1500 | Total |
| Tongue and mouth (C01–C06) | Cases | 3 | 12 | 5 | 11 | 1 | 6 | 0 | 38 | 0.045 | 0.50 |
| RR                        | 1.00 | 1.30 | 3.10 | 3.12 | 0.43 | 6.15 | — | — | — | — | — | — |
| Pharynx (C09–C14) | Cases | 6 | 16 | 2 | 10 | 12 | 6 | 1 | 53 | 0.163 | 0.12 |
| RR                        | 1.00 | 0.97 | 0.54 | 1.27 | 2.85 | 3.33 | 1.18 | — | — | — | — | — |
| Oesophagus (C15) | Cases | 19 | 38 | 7 | 37 | 15 | 6 | 3 | 125 | 0.025 | 0.08 |
| RR                        | 1.00 | 0.97 | 0.76 | 1.31 | 0.82 | 0.69 | 0.68 | — | — | — | — | — |
| Stomach (C16) | Cases | 76 | 143 | 35 | 141 | 95 | 62 | 38 | 590 | 0.021 | 0.04 |
| RR                        | 1.00 | 1.21 | 1.42 | 1.34 | 1.16 | 1.51 | 1.77 | — | — | — | — | — |
| Colon (C17–C18) | Cases | 51 | 74 | 13 | 66 | 41 | 44 | 10 | 299 | 0.017 | 0.26 |
| RR                        | 1.00 | 0.79 | 0.63 | 0.84 | 0.71 | 1.58 | 0.71 | — | — | — | — | — |
| Rectum (C19–C21) | Cases | 39 | 55 | 16 | 53 | 39 | 22 | 17 | 241 | 0.028 | 0.13 |
| RR                        | 1.00 | 0.77 | 0.97 | 0.91 | 0.89 | 1.03 | 1.58 | — | — | — | — | — |
| Liver (C22) | Cases | 25 | 35 | 12 | 24 | 34 | 21 | 7 | 158 | 0.044 | 0.09 |
| RR                        | 1.00 | 0.70 | 1.10 | 0.59 | 1.18 | 1.52 | 1.02 | — | — | — | — | — |
| Gallbladder (C23–C24) | Cases | 15 | 17 | 3 | 19 | 12 | 9 | 6 | 81 | 0.021 | 0.46 |
| RR                        | 1.00 | 0.70 | 0.57 | 0.85 | 0.70 | 1.05 | 1.37 | — | — | — | — | — |
| Pancreas (C25) | Cases | 38 | 60 | 11 | 51 | 43 | 16 | 9 | 228 | 0.001 | >0.5 |
| RR                        | 1.00 | 0.81 | 0.70 | 0.93 | 1.04 | 0.78 | 0.86 | — | — | — | — | — |
| Larynx (C32) | Cases | 13 | 15 | 2 | 16 | 20 | 8 | 1 | 75 | 0.021 | 0.49 |
| RR                        | 1.00 | 0.66 | 0.40 | 0.92 | 1.51 | 1.23 | 0.29 | — | — | — | — | — |
| Prostate (C61) | Cases | 50 | 60 | 12 | 62 | 42 | 20 | 17 | 263 | 0.000 | >0.5 |
| RR                        | 1.00 | 0.85 | 0.81 | 0.88 | 0.75 | 0.71 | 1.20 | — | — | — | — | — |
| Kidney (C64–C66) | Cases | 26 | 44 | 12 | 42 | 23 | 13 | 11 | 171 | 0.017 | 0.39 |
| RR                        | 1.00 | 0.83 | 0.96 | 1.01 | 0.79 | 0.92 | 1.80 | — | — | — | — | — |
| Bladder (C67–C68) | Cases | 22 | 34 | 10 | 42 | 39 | 22 | 8 | 177 | 0.020 | 0.28 |
| RR                        | 1.00 | 1.03 | 1.47 | 1.26 | 1.44 | 1.60 | 1.15 | — | — | — | — | — |
| Brain and others (C70–C72) | Cases | 15 | 41 | 9 | 18 | 22 | 10 | 0 | 115 | 0.018 | 0.27 |
| RR                        | 1.00 | 1.32 | 1.33 | 0.85 | 1.56 | 1.52 | — | — | — | — | — | — |
| Non-Hodgkin’s disease (C82–C85 and C91.4) | Cases | 14 | 22 | 8 | 18 | 13 | 9 | 3 | 87 | 0.032 | 0.35 |
| RR                        | 1.00 | 0.71 | 1.34 | 1.00 | 1.05 | 1.54 | 1.05 | — | — | — | — | — |
| Myeloma (C90) | Cases | 13 | 11 | 2 | 14 | 7 | 7 | 1 | 55 | 0.007 | >0.5 |
| RR                        | 1.00 | 0.40 | 0.37 | 0.82 | 0.56 | 1.22 | 0.34 | — | — | — | — | — |
| Leukaemia (C91–C95, excluding C91.4) | Cases | 22 | 29 | 5 | 17 | 14 | 4 | 127 | 0.006 | >0.5 |
| RR                        | 1.00 | 0.75 | 0.57 | 1.13 | 0.75 | 1.26 | 0.71 | — | — | — | — | — |
| Other and unspecified (C91–C95, excluding C91.4) | Cases | 75 | 137 | 24 | 93 | 74 | 31 | 23 | 457 | 0.009 | 0.44 |
| RR                        | 1.00 | 0.96 | 0.84 | 0.90 | 1.01 | 0.86 | 1.29 | — | — | — | — | — |
| All non-lung cancers | Cases | 522 | 843 | 188 | 753 | 549 | 326 | 159 | 3340 | 0.014 | <0.001 |
| RR                        | 1.00 | 0.89 | 0.94 | 1.00 | 0.99 | 1.20 | 1.16 | — | — | — | — | — |
| Person-years at risk | 0-year lag | 256,264 | 781,696 | 102,726 | 290,243 | 191,867 | 92,371 | 47,019 | 1,762,208 | — | — | — |
| 5-year lag | 363,845 | 693,782 | 95,229 | 287,756 | 191,037 | 86,643 | 43,913 | 1,762,208 | — | — | — | — |

ERR/WLM = excess relative risk per working level months. RR = relative risk. *Statistically significant (P < 0.05).
exposure to external γ-radiation, LRN, arsenic and dust led to some decrease in the ERR/WLM. Confounding from other factors such as alcohol consumption cannot be ruled out.

**Stomach** A significantly elevated SMR for stomach cancer (n = 590, O/E = 1.15; 95% CI: 1.06–1.25) was observed here, as in other studies of radon-exposed miners (Kusiak et al, 1993; Darby et al, 1995a), and among coal miners (Rockette, 1977). Although not fully understood, it could be related to dust exposure (Cocco et al, 1996). In the pooled study, an elevated SMR (n = 217; SMR = 1.33; 95% CI: 1.16–1.52) was found with no exposure–response relationship (Darby et al, 1995a), whereas in our study, the risk increased significantly with increasing cumulative radon exposure (ERR/WLM = 0.022%). The highest exposure category (1500 WLM or more) was associated with a 1.8-fold (95% CI: 1.06–2.48) significantly higher risk of death compared with the reference category of 0 WLM. Adjustment for each of the five confounders, however, reduced the ERR/WLM by a factor of approximately 2, leading to insignificant values. Thus, part of the proportionate increase in risk because of radon might be explained by confounding.

**Pharynx** A significant deficit of cancers of the tongue, mouth, salivary gland and pharynx combined (n = 99, SMR = 0.8; 95% CI: 0.65–0.97) may be a chance finding because of multiple testing. There was a constant, but not significant, increase in pharyngeal cancer risk with increasing cumulative radon exposure (n = 53, ERR/WLM = 0.16%; 95% CI: −0.045 to 0.37%). It can be noted that this value was nearly as high as for lung cancer (ERR/WLM = 0.20%), but no such relationship was reported in other studies on miners, although the number of pharyngeal cases was small (Tomasek et al, 1993; Darby et al, 1995a). Additional adjustment for the five possible confounders only led to a small reduction of the ERR/WLM. Some studies have provided estimates for organ doses after inhalation of radon and its progeny separately for the extrathoracic airways, and have reported a pharyngeal dose that was nearly as high as the lung dose (Kendall and Smith 2002; Jacobi and Roth, 1995).

**Larynx** The combined 11 studies on miners showed a 1.21-fold non-significantly increased SMR for larynx cancer that was not related to cumulative radon exposure (Darby et al, 1995a), but there were only 38 cases. In the first follow-up of the French uranium miners’ study (1946–1985), a significantly increased SMR of 2.35 was observed on the basis of 17 cases (Tirmarche et al, 1993), which became insignificant after extension of the follow-up period to 1999 (SMR = 1.24, n = 29) (Vacquier et al, 2008). The SMR in this study (n = 75; SMR = 1.18; 95% CI: 0.93–1.48) is comparable with the findings of the pooled study (Darby et al, 1995a). ERR/WLM was elevated, but not significantly. Adjustment for the five potential confounders led to a substantial decrease in the observed ERR/WLM.

**Kidney** Animal experiments suggest an increased mortality of kidney cancer after inhalation of radon (Masse et al, 1992), but none of the miners’ studies found any such excess (Tomasek et al, 1993; Darby et al, 1995a), apart from the French study (n = 20, SMR = 2.0; 95% CI: 1.22–3.09) (Vacquier et al, 2008). Moreover, none of these studies observed a trend with cumulative radon exposure. The same holds true in our data, there being no excess (n = 162, SMR = 0.91) or an exposure–response relationship.
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Leukaemia In our study, no association between cumulative radon exposure and leukaemia is found, or with CLL, non-CLL or AML, consistent with earlier studies (Tomasek et al, 1993; Darby et al, 1995a; Laurier et al, 2001; Vacquier et al, 2008), including a recent large case-control study with 377 leukaemia cases among former Wismut employees (Moehner et al, 2006). In contrast, Rericha et al (2006) noted, in a Czech uranium miner case-control study, a significantly increased relative risk of 1.75 (95% CI: 1.10–1.75) for leukaemia incidence in the highest quintile of cumulative radon exposure (>100 WLM) compared with the lowest (<3 WLM). However, there was a very high correlation in the mines, between radon exposure and exposure to γ-radiation, which could have introduced confounding bias.

Strengths and limitations

The major strengths of our study are the large cohort size, the large number of extrapulmonary cancers, the long follow-up period, the wide range of radon exposures and particularly the information available on other exposures such as arsenic, fine dust, silica, external γ-radiation and LRN. These advantages allowed the independent replication of the analysis of the 11 miners’ cohort studies (Darby et al, 1995a), which may have suffered from heterogeneity problems. The potential limitations of this study include the accuracy of the underlying causes of death on death certificates, missing causes of death, exposure misclassification particularly in the early years of mining activities as well as missing information on other potential confounders such as alcohol consumption, smoking, occupational exposure to diesel exhaust or asbestos. Moreover, despite the large number of cancer cases overall, there is a low statistical power with respect to certain sites, and multiple testing could have led to some spurious findings.

Confounding

Within a nested case-control study of lung cancer in the Wismut cohort, information on smoking was collected from miners, their relatives and the medical Wismut archive. Most of the former Wismut employees had been smokers. Overall, the low correlation between smoking and cumulative radon exposure makes smoking an unlikely major confounder. It is known that Wismut employees in the early years had a relatively high alcohol consumption compared with the male general population. For approximately 5% of the deceased cohort members, alcohol abuse was noted on the death certificate. This rough surrogate for alcohol consumption was slightly negatively correlated with cumulative radon exposure.

Exposure misclassification

Inevitably, exposures in the very early years are associated with considerable uncertainty. To obtain some insight into potential bias by misclassification, the cumulated radon exposure was separated into two components according to other studies investigating the effect of the quality of exposure (Tomasek et al, 2008; Vacquier et al, 2008), one risk estimate for the period 1946–1954, the years with retrospectively estimated radon concentrations and the other for when the JEM was based on measurements in the shafts. As, for all non-lung cancers combined, there was only a non-significant difference in the estimates for these two periods, a major bias through misclassification of exposure is unlikely, but cannot be excluded. Another potential limitation is the use of the exposure to radiation instead of the actual organ dose. Recently, it has been suggested that several factors such as physical activity, ventilation in the mines, dry or wet drilling may influence the individual doses (Marsh et al, 2008). Work on these dose calculations is currently in progress within the European collaborative research project ALPHA-RISK (European Commission, 2006), which will also provide a method for calculating the dose to the various organs from combined exposure to radon and its progeny, LRN and external γ-radiation.

CONCLUSION

Some evidence of a very small radon-related risk of extrapulmonary cancers was found, compatible with dosimetric calculations for organ doses. However, the possibility of non-causal results because of chance and confounding cannot be ruled out.

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