Effects of occupational exposure to respirable quartz dust on acute myocardial infarction

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ABSTRACT

Objectives The aim of this study is to investigate the effects of occupational exposure to respirable quartz (RQ) on first acute myocardial infarction (AMI). RQ causes pulmonary diseases like silicosis and has also been linked to cardiovascular diseases. Inflammation is hypothesised as the underlying pathway.

Methods We performed a 1:3 matched case–control study nested in a cohort of male uranium miners. We included cases (identified from hospital records and validated according to WHO criteria) who had suffered their first AMI while still employed and <65 years of age. Controls were matched by date of birth and Wismut recruitment era. RQ exposure was derived from a job-exposure matrix. We performed a conditional logistic regression adjusted for smoking, metabolic syndrome and baseline erythrocyte sedimentation rate. Subgroups by date of birth and Wismut recruitment era were analysed to minimise the impact of pre-exposures.

Results The study base comprised 292 matched sets. The cumulative exposure ranged from 0 to 38.9 mg/m²-years RQ. The adjusted OR of the highest RQ tertile (>14.62 mg/m²-years) was 1.27 (95% CI 0.82 to 1.98). However, for miners born after 1928 and hired in the earliest recruitment era (1946–1954), a significantly elevated risk was seen in the highest RQ tertile (OR=6.47 [95% CI 1.33 to 31.5]); 50 matched sets).

Conclusions An impact of quartz dust on first AMI was observed only in a small subgroup that had virtually no pre-exposure to RQ. Further studies on the basis of complete occupational history are required to substantiate this finding.

INTRODUCTION

Respirable quartz (RQ) has long been known to cause silicosis. In 1997, the International Agency for Research on Cancer (IARC) determined that crystalline silica causes lung cancer. Both silicosis and lung cancer are believed to result from the strong inflammatory response that silica evokes in the lung. Furthermore, other conditions like chronic obstructive pulmonary disease1 and cardiovascular disease2 have been associated with RQ. The exact biological mechanisms linking exposure to RQ with cardiovascular disease are still unclear.

Potential mechanisms discussed in the literature include systemic inflammation following primary inflammation of the lungs. There is growing evidence that inflammation is a key regulatory process in the pathophysiology of atherosclerosis. A cohort study demonstrated that the erythrocyte sedimentation rate (ESR) is a long-term independent predictor of coronary heart disease.7 Taken together, RQ seems to be a first-rate candidate to cause inflammation-mediated acute myocardial infarction (AMI). However, the number of studies investigating the impact of occupational RQ dust on the AMI-risk yields inconsistent results.

Some cohort studies report elevated risks of coronary heart disease mortality by RQ,3,4 yet this has not been confirmed by other cohort studies.5 The same applies to case–control studies.6 The number of studies investigating the impact of occupational RQ dust on AMI is still unclear.

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of death, this definition of the endpoint does not differentiate between first and recurrent AMI events and obscures the time of onset. The time lag between first AMI and death may be considerable, thus shifting the focus of the study to older age. The temporal reference to occupational dust may lose precision and, moreover, be biased by job changes due to earlier cardiovascular events (eg, angina pectoris). Therefore, we investigated the impact of RQ on the first AMI in a nested case–control study.

**METHODS**

**Study population**

This study is nested in the German Uranium Miners cohort study (Wismut cohort) run by the Federal Office for Radiation Protection (BfS). The cohort comprises a sample of approximately 60,000 employees who worked in uranium ore mining between 1946 and 1990. The cohort was stratified according to the period of hiring (1946–1954, 1955–1970, 1971–1989, subsequently denoted as subcohorts A, B and C). These strata represent different mining conditions in terms of exposure to radiation and dust, with different sampling probabilities for the strata, too.

In parallel, the Federal Institute for Occupational Safety and Health (BAuA) runs the health data archives Wismut (GDAW), which provide 12,000 linear metres of medical records and radiography of former uranium miners, including medical records on health surveillance, hospital discharge diagnosis and occupational diseases. All hospitalisations of the cohort were digitised. This file comprises 114,000 hospitalisations in nearly 28,000 ever hospitalised patients. The hospital discharge diagnoses are coded in ICD-10. We searched for ICD-10: I21, I22, I25.2, I29.9+ plaintext and thus identified 676 potential first AMI cases. To validate these, we reviewed inpatient and outpatient medical records.

We applied the 1979 WHO definition of myocardial infarction, which requires at least two out of three criteria (clinical symptoms, ECG evaluation and laboratory values) to be fulfilled. We then excluded (1) 6 cases where no suitable controls were available, (2) 11 cases lacking exposure data, (3) 175 cases with AMI dated after leaving the company. The latter restriction was necessary because the information on exposure and outcome can be assumed to be complete only for the period of Wismut-employment. Controls were individually matched at a ratio of 3:1 by date of birth (± 1 year) and period of hiring. The event date of the case defines the index date for the respective case–control set. At the index date, all controls were as old as the corresponding case, had been employed at the Wismut company, had benefited from the company’s health system and were free of AMI.

**Figure 1** Flowchart of the inclusion process of cases. AMI, acute myocardial infarction.

**Classic risk factors and markers of systemic inflammation**

Information on classic risk factors of atherosclerosis, that is, smoking habits, blood pressure, cholesterol and blood glucose level, and on ESR as a marker of systemic inflammation were collected from medical records covering the period of commencement of employment until the index date. The presence of hypertension, hypercholesterolaemia and diabetes was presumed if either a medical diagnosis was given as a plain text, or a respective medication or at least three elevated laboratory values were ascertained (thresholds: hypertension 160/100 mm Hg, hypercholesterolaemia ≥6.2 mmol/L, diabetes ≥7.0 mmol/L). The diagnoses and treatment decisions were based on the state of knowledge during the study period 1950–1989 applying thresholds at that time. Therefore, we did not choose current thresholds for the analysis. Information on classic risk factors 5 years prior to index date were omitted to avoid a lead time bias, that is, the inflated chance to detect a case’s risk factors because of early cardiovascular symptoms. We defined the metabolic syndrome by the presence of ≥3 out of 4 criteria (obesity, hypertension, diabetes and hypercholesterolaemia), approximating the International Diabetes Federation (IDF) metabolic syndrome consensus definition. ESR was considered elevated for a 1-hour value >10 mm and a 2-hour value >20 mm.

**Exposure assessment**

The BfS provided exposure measures per person and year that combine the job-exposure matrix with the individual job history. The cumulative RQ exposure is given in mg/m³-years, which is defined as an exposure to 1 mg/m³ over 220 shifts of 8 hours each. The individual cumulative RQ exposure was calculated omitting the 5 years preceding the index date. We thus aimed to avoid a healthy-worker effect, which can arise if early cardiovascular symptoms cause a change from an exposed to an unexposed job. The extent of exposure prior to the Wismut employment cannot be quantified, even if some qualitative binary information on pre-exposure is available as a part of the personnel index card accompanying the preventive medical examinations. Our first concern was pre-exposure during military service, internment (for example, as a prisoner of war), or (pre-war) coal mining. To manage this, we created a further binary categorisation whether a subject was older or younger than 16 years at the end of World War II (born before or after 8 May 1929, subsequently denoted as early and late birth cohorts). Our second concern was that, in general, every employee who entered the company beyond young adulthood may have gathered occupational pre-exposure beforehand. The Wismut
company hired experienced coal miners in the 1960s (subcohort B) to establish a permanent staff. With regard to the study objective, pre-exposure is less harmful if it is small compared with relatively high measured exposure. Hence, we investigated the six subgroups, defined by their birth cohort (early, late) and period of hiring (A, B and C), to find out a subgroup with a chance of occupational pre-exposure to RQ as low as possible.

**Statistical analysis**

To account for different time-related changes a tight matching was used, taking into account date of birth and period of hiring. Hence, a matched analysis was required. ORs and two-sided 95% CIs were calculated by conditional logistic regression. Cumulative exposure was categorised into tertiles to account for non-linearity in a possible exposure–response relationship. The model fit was assessed by means of a corresponding likelihood-ratio test, as well as by Akaike’s information criterion (AIC). All statistical analyses were performed using the STATA software package V.15. Smoking, the baseline ESR and the metabolic syndrome (combined from hypertension, diabetes, hypercholesterolaemia and obesity) were considered as potential confounders.

To better control the impact of the confounding variables on the RQ-related risk estimates, we applied three models and calculated: the crude OR, the OR adjusted for the potential confounders smoking and baseline ESR, and finally the OR with additional adjustment for the metabolic syndrome as a suspected mediator.

**Table 1** Description of the study population

| Date of birth | Before 08.05.1929 (early birth cohorts) | After 08.05.1929 (late birth cohorts) | Total |
|---------------|----------------------------------------|-------------------------------------|-------|
|               | Cases | Controls | Cases | Controls | Cases | Controls |
| n             | 182   | 542      | 110   | 330      | 292   | 872      |
| Year of birth, mean (SD) | 1919 (7.3) | 1919 (7.3) | 1935 (5.5) | 1935 (5.5) | 1925 (10.0) | 1925 (10.0) |
| Year of hiring, mean (SD) | 1950 (4.3) | 1950 (4.3) | 1957 (8.0) | 1957 (8.0) | 1953 (7.0) | 1953 (6.6) |
| Age at hire, mean (SD) | 31.1 (8.8) | 31.4 (8.7) | 22.0 (6.5) | 22.0 (5.7) | 27.7 (9.1) | 27.9 (8.9) |
| Age at index date*, mean (SD) | 55.2 (5.8) | 55.3 (5.7) | 47.9 (6.6) | 47.9 (6.6) | 52.5 (7.1) | 52.5 (7.0) |
| Time since hire, mean (SD) | 24.1 (8.7) | 23.9 (8.7) | 25.9 (9.0) | 26.0 (8.5) | 24.8 (8.9) | 24.7 (8.7) |
| Last smoking status, n (%) | | | | | | |
| Non-smoker | 14 (7.7) | 116 (21.4) | 6 (5.5) | 81 (24.5) | 20 (6.8) | 197 (22.6) |
| Ex-smoker | 28 (15.4) | 68 (12.5) | 13 (11.8) | 48 (14.5) | 41 (14.0) | 116 (13.3) |
| Smoker | 123 (67.6) | 245 (45.2) | 89 (80.9) | 164 (49.7) | 212 (72.6) | 409 (46.9) |
| Unknown | 17 (9.3) | 113 (20.8) | 2 (1.8) | 37 (11.2) | 19 (6.5) | 150 (17.2) |
| Diabetes mellitus†, n (%) | 15 (8.2) | 7 (1.3) | 12 (10.9) | 8 (2.4) | 27 (9.3) | 15 (1.7) |
| Hypertension†, n (%) | 57 (31.3) | 108 (19.9) | 39 (35.5) | 83 (25.2) | 96 (32.9) | 191 (21.9) |
| Hypercholesterolaemia†, n (%) | 3 (1.6) | 8 (1.5) | 13 (11.8) | 5 (1.5) | 16 (5.5) | 13 (1.5) |
| Obesity‡, n (%) | 17 (9.3) | 59 (10.9) | 21 (19.1) | 44 (13.3) | 38 (13.0) | 103 (11.8) |
| Erythrocyte sedimentation rate, earliest/baseline measure | | | | | | |
| Elevated §, n (%) | 43 (24.9) | 86 (16.2) | 21 (19.1) | 30 (9.1) | 64 (21.9) | 116 (13.3) |
| 1 hour, mean (SD) | 8.6 (12.0) | 6.8 (8.6) | 7.3 (10.1) | 5.9 (11.1) | 8.1 (11.3) | 6.5 (9.6) |
| 2 hours, mean (SD) | 17.8 (18.6) | 14.8 (14.5) | 16.3 (17.0) | 12.6 (15.0) | 17.2 (18.0) | 14.0 (14.7) |
| Metabolic syndrome¶, n (%) | 7 (3.8) | 5 (0.9) | 7 (6.4) | 6 (1.8) | 14 (4.8) | 11 (1.3) |
| Cumulative RQ**, (mg/m³-years), mean (SD) | 12.9 (8.9) | 13.18 (9.77) | 9.12 (8.68) | 8.42 (8.23) | 11.5 (9.00) | 11.38 (9.50) |

Results

Six hundred seventy six potential cases of AMI were identified from hospital discharge diagnoses, resulting in 484 valid cases according to the 1979 WHO definition of myocardial infarction. Reviewing the complete medical records ensured that only primary events of myocardial infarction were captured.

Exclusion of 192 cases for various reasons resulted in 292 cases for the main analysis (figure 1). These were matched with 872 controls, that is, most cases matched 3 controls. The date of birth of the controls differed by 4.4 days on average (maximum=246 days) from the date of birth of the respective case. Table 1 shows the main characteristics of the study population by the two birth-cohort strata. The study population was born between 1902 and 1960 and commenced employment between 1946 and 1986. The subgroups differ considerably from each other. Miners from subcohort A had on average more than 35-fold cumulative exposure in comparison to miners from subcohort C (table 2). The age at hire also differs considerably between the six subgroups formed by the birth cohorts and the period of hiring. The lowest average age at hire was observed for the subgroup of the late birth cohort from subcohort A. Hence, the impact of relevant pre-exposures should be negligible in this subgroup. Moreover, the period between hire and index date covers on average more than 30 years in this subgroup and is therefore significantly longer in comparison to all other subgroups.

RQ exposure strongly depends on calendar time (figure 2). Within the study population, the median exposure concentration peaked in 1952 with 1.6 mg/m³. It then decreased to 0.3 in 1963 and further to 0.03 in 1970.
In view of the impact of potential pre-exposure, risk analyses were performed based on the entire database as well as for the late birth cohorts and, additionally, for the late birth cohorts restricted to subcohort A, due to their low chance of considerable occupational pre-exposure to RQ.

Univariate risk estimates (online supplementary table S1) indicate strong associations of AMI with smoking and with the infrequent risk factors diabetes, the metabolic syndrome and hypercholesterolaemia.

We applied the categorised cumulative RQ exposure with the lowest RQ tertile as referent in the main analysis, where the tertiles were calculated based on all AMI cases. The analysis of the entire database does not reveal a clear relationship between RQ exposure and risk of AMI, even if the OR for the middle tertile is significantly elevated (table 3). Although the analysis restricted to the late birth cohort suggests such a relationship, the significance threshold is not exceeded even in the highest tertile. The analysis of the late birth cohorts from subcohort A, however, yields a significant increase in the risk of AMI with OR=6.47 (95%CI: 1.33 to 31.5) in the highest tertile. The analysis of the late birth cohorts from subcohort A in the fully adjusted model. A much better model fit was achieved by a restricted to the late birth cohort suggests such a relationship, the significance threshold is not exceeded even in the highest tertile. The analysis of the late birth cohorts from subcohort A, however, suggests such a relationship, the significance threshold is not exceeded even in the highest tertile.

We also included the exposure as a linear term instead of tertiles. While in the entire database an increased risk could not be observed (OR=1.00 [95% CI 0.98 to 1.02] per 1 mg/m³-year), it was OR=1.04 (95% CI 0.99 to 1.09; p=0.10) for the subgroup of late birth cohorts from subcohort A in the fully adjusted model. A much better model fit was achieved by a
The likelihood-ratio test confirms the significant impact of RQ on the AMI-risk, both for the model based on tertiles and for the model based on the log-transformed exposure. The comparison of both models by means of AIC is in favour of the categorical model. The ORs of the exposure were similar throughout all three models in the complete database as well as in the subgroups regardless of the respective covariates.

### DISCUSSION

Given the complete study base, there is no statistically significant impact of occupational quartz dust on the first AMI. In contrast, the analysis restricted to the subgroup of late birth cohorts from subcohort A showed a significant risk increase. The decisive argument to highlight this subgroup is that it is virtually unaffected by unmeasured pre-exposure to RQ: Its subjects commenced employment at young age (19 years on average) in a period of very high occupational exposures and they stayed with the company for many years (31 years on average). However, this subgroup comprises only 50 matched sets, ie, 17% of the study population. Hence, our study adds to the still small body of evidence for an exposure–response relationship between RQ and AMI risk.

A major strength of our study is that all potential study subjects were reviewed using medical records from the comprehensive GDAW archives.

This procedure largely ensures that only first events, but no reinfections, are included as cases in the study whereas potential controls with hints at previous AMI are excluded. In contrast, in mortality-based studies only the fatal AMI is observed. Thus, possible previous non-fatal AMI events, requiring a change of job with a respective decrease of exposure, could lead to a healthy-worker survivor bias. In order to minimise the risk of such a bias, we lagged the exposures by 5 years. The use of unlagged cumulative exposure leads virtually to the same results, since the cumulative exposure in the last 5 years is for almost all subjects negligible in comparison to the previous exposure (data not shown).

Reports of preventive medical examinations provided also important data on non-occupational cardiovascular risk factors. We considered these risk factors as potential confounders or mediators in our models (tables 1 and 3, online supplementary tables S1, S2). We omitted the information on risk factors that only turned out within the 5 years ahead of the index date. However, the results with and without this lag time did not differ much (online supplementary table S2).

Interestingly, the RQ-related ORs showed similarities, regardless of the covariates introduced in the respective model (table 3). This suggests that smoking, baseline ESR and the metabolic syndrome, although very predictive for AMI, do not generally confound the estimation. The metabolic syndrome had been suspected to mediate the effect of RQ on AMI, but it did not, as the introduction of this covariate did not lower the RQ-related ORs, neither in the analysis of the complete database nor in the subgroups. Thus, the metabolic syndrome seems to be an independent risk factor for AMI.

We are aware of some weaknesses of our study. Underdetection of cases cannot be ruled out. This weakness is unavoidable as there is neither a national AMI registry nor a hospital discharge registry for Germany. Another source of underdetection of cases in this study could be the absence of emergency medical services until the mid-1960s. Until then, the protection of cases in this study might have been much higher. Another weakness of our study is that RQ is not the only hazard uranium miners are exposed to. We had to ask whether a possible impact of RQ is not in truth an impact of ionising radiation. We thus used the study design to alternatively explore the impact of radon, gamma radiation and long-lived radionuclides on AMI. Despite strong correlations of all hazards mentioned, but in contrast, the analysis restricted to the subgroup of late birth cohorts only* from subcohort A only*

| RQ (mg/m³-years) | Entire database | Late birth cohorts only* | Late birth cohorts only* from sub-coh ort A only* |
|-----------------|----------------|--------------------------|-----------------------------------------------|
| Inclusion of RQ as | Cases Controls OR (95% CI) | Cases Controls OR (95% CI) | Cases Controls OR (95% CI) |
| Tertiles† | | | |
| <5.74 | 97 330 1.00 (referent) | 53 166 1.00 (referent) | 2 29 1.00 (referent) |
| 5.74 to <14.62 | 97 247 1.52 (1.02 to 2.25) | 27 92 1.03 (0.53 to 1.99) | 20 53 5.42 (1.19 to 24.8) |
| ≥14.62 | 98 295 1.32 (0.87 to 1.99) | 30 72 1.59 (0.74 to 3.42) | 28 68 6.36 (1.37 to 29.6) |
| Tertiles‡ | | | |
| <5.74 | 97 330 1.00 (referent) | 53 166 1.00 (referent) | 2 29 1.00 (referent) |
| 5.74 to <14.62 | 97 247 1.65 (1.05 to 2.51) | 27 92 1.19 (0.56 to 2.51) | 20 53 6.53 (1.35 to 31.5) |
| ≥14.62 | 98 295 1.25 (0.80 to 1.93) | 30 72 1.58 (0.67 to 3.69) | 28 68 6.55 (1.35 to 31.8) |
| Tertiles§ | | | |
| <5.74 | 97 330 1.00 (referent) | 53 166 1.00 (referent) | 2 29 1.00 (referent) |
| 5.74 to <14.62 | 97 247 1.74 (1.14 to 2.65) | 27 92 1.26 (0.59 to 2.68) | 20 53 6.46 (1.34 to 31.3) |
| ≥14.62 | 98 295 1.27 (0.82 to 1.98) | 30 72 1.64 (0.69 to 3.88) | 28 68 6.47 (1.33 to 31.5) |
| Linear term§ | | | |
| RQ | 292 872 1.00 (0.98 to 1.02) | 110 330 1.03 (0.99 to 1.07) | 50 150 1.04 (0.99 to 1.09) |
| Ln(1+RQ) | 292 872 1.11 (0.90 to 1.37) | 110 330 1.23 (0.82 to 1.83) | 50 150 1.97 (1.05 to 3.71) |

*Date of birth after 8 May 1929.
†Crude analysis.
‡With smoking and first erythrocyte sedimentation rate.
§With additional adjustment for metabolic syndrome.

Table 3 ORs for the first acute myocardial infarction by cumulative exposure to respirable quartz (RQ)
cohort studies have been discussed as an explanation of the inconsistencies in literature. \(^9\)

The present study reflects an exposure scenario in uranium ore mining based on data from 1946 to 1990 with very high annual RQ exposure, especially in the first decade (figure 2). Permissible limit values for RQ dust have been reduced substantially since. But silicosis is still among the 10 most frequent recognised occupational diseases in Germany. \(^23\) Even today the hazard of high occupational exposure to RQ is not averted. \(^24\) \(^25\) \(^26\) \(^27\)

To conclude, although a significant risk increase was only seen in a small subgroup of our study, a link between RQ exposure and AMI could not be ruled out. Further investigations on this topic are desirable. Such a study requires a thorough validation of the outcome based on clinical data, as well as reliable exposure information on the subjects' complete occupational history.

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**Contributors** JG and MM conceived the research question. JG checked the medical records of all study subjects and verified all AMI diagnoses. DP supervised the data collection, programmed the study database and carried out quality controls. MM performed the statistical analysis. JG wrote the first draft of the manuscript. All authors revised it critically and approved the final version. MM is responsible for the overall content as guarantor.

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**REFERENCES**

1. Steenland K, Ward E. Silica: a lung carcinogen. *CA Cancer J Clin* 2014;64:63–9.

2. Möhrer M, Kersten N, Gellissen J. Chronic obstructive pulmonary disease and longitudinal changes in pulmonary function due to occupational exposure to respirable quartz. *Occup Environ Med* 2013;70:9–14.

3. Weiner J, Barlov L, Sjögren B. Ischemic heart disease mortality among miners and other potentially silica-exposed workers. *Am J Ind Med* 2007;50:403–8.

4. Fang SC, Cassidy A, Christiani DC. A systematic review of occupational exposure to particulate matter and cardiovascular disease. *Int J Environ Res Public Health* 2010;7:1773–806.

5. Libby P, Ridker PM, Hansson GK, et al. Inflammation in atherosclerosis: from pathophysiology to practice. *J Am Coll Cardiol* 2009;54:2129–38.

6. Hansson GK. Inflammation, atherosclerosis, and coronary artery disease. *N Engl J Med* 2005;352:1685–95.

7. Andresdottir MB, Sigfusson N, Sigvaldason H, et al. Erythrocyte sedimentation rate, an independent predictor of coronary heart disease in men and women: The Reykjavik Study. *Am J Epidemiol* 2003;158:844–51.

8. Torén K, Bergdahl IA, Nilsson T, et al. Occupational exposure to particulate air pollution and mortality due to ischaemic heart disease and cerebrovascular disease. *Occup Environ Med* 2007;64:515–9.

9. Fan C, Graff P, Vilhberg P, et al. Silica exposure increases the risk of stroke but not myocardial infarction: A retrospective cohort study. *PloS One* 2018;13:e0192840.

10. Sjögren B. Occupational exposure to dust: inflammation and ischaemic heart disease. *Occup Environ Med* 1997;54:466–9.

11. Hammar N, Alfredsson L, Smedberg M, et al. Differences in the incidence of myocardial infarction among occupational groups. *Scand J Work Environ Health* 1992;18:178–85.

12. Wyndham CH, Bezuidenhout BN, Greenacre MJ, et al. Mortality of middle aged white South African gold miners. *Br J Ind Med* 1986;43:677–84.

13. Reid PI, Sluis-Cremer GK. Mortality of white South African gold miners. *Occup Environ Med* 1996;53:11–16.

14. Kreuzer M, Grosche B, Dufey F, et al. The German Uranium Miners Cohort Study (Wismut cohort), 1946-2003. *Oberschlesien: Bundesamt für Strahlenschutz* 2011:50.

15. Grosche B, Kreuzer M, Kreisheimer M, et al. Lung cancer risk among German male uranium miners: a cohort study, 1946-1998. *Br J Cancer* 2006;95:1280–7.

16. IDF. The IDF consensus worldwide definition of the metabolic syndrome. Brussels: IDF; 2006:24.

17. Alberti KG, Zimmet P, Shaw J. Metabolic syndrome--a new world-wide definition. *A Consensus Statement from the International Diabetes Federation. Diabet Med* 2002;23:469–80.

18. Dahmann D, Bauer HD, Stoyke G. Retrospective exposure assessment for respirable and inhalable dust, crystalline silica and arsenic in the former German uranium mines of SAG/SAG-Wismut. *Int Arch Occup Environ Health* 2008;81:949–58.

19. Runge W, ed. *Chronik der Wismut. Chemnitz: Wismut GmbH*, 1999.

20. Schisselmann JJ. Case-control studies. Design, conduct and analysis. New York: Oxford Univ Press, 1982:368.

21. StataCorp. *Stata: Release 15. Statistical Software*. College Station, TX: StataCorp LP; 2017.

22. Mente A, Yusuf S, Islam S, et al. Metabolic syndrome and risk of acute myocardial infarction: a case-control study of 26,903 subjects from 52 countries. *J Am Coll Cardiol* 2010;55:2390–8.

23. Anand SS, Islam S, Rosengren A, et al. Risk factors for myocardial infarction in women and men: insights from the INTERHEART study. *Eur Heart J* 2008;29:932–40.

24. Yusuf S, Hawken S, Ounpuu S, et al. Effect of potentially modifiable risk factors associated with myocardial infarction in 52 countries (the INTERHEART study): case-control study. *Lancet* 2004;364:937–52.

25. Darby SC, Ewertz M, McGale P, et al. Risk of ischemic heart disease in women after radiotherapy for breast cancer. *N Engl J Med* 2013;368:987–98.

26. Kreuzer M, Kreisheimer M, Kandel M, et al. Mortality from cardiovascular diseases in the German uranium miners cohort study, 1946-1998. *Radiat Environ Biophys* 2006;45:159–66.

27. DGU/BBK-Monitoring-Bericht Berufskrankheiten im Jahr 2016. \(^2016\) 2018.

28. Laney AS, Attfield MD. Coal workers’ pneumoconiosis and progressive massive fibrosis are increasingly more prevalent among workers in small underground coal mines in the United States. *Occup Environ Med* 2010;67:428–31.

29. Perret JL, Plush B, Lachapelle P, et al. Coal mine dust lung disease in the modern era. *Respirology* 2017;22:662–70.

Gellissen J, et al., *Occup Environ Med* 2019; 76:370–375. doi:10.1136/oemed-2018-105540