PULMONARY DYSFUNCTION IN INDIUM TIN OXIDE EXPOSED WORKERS

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Aim To investigate the relationship between indium exposure and effects markers among indium tin oxide (ITO) manufacturing workers without job change.

Methods We enrolled 179 male workers from ITO target manufacturing and recycling factories in Taiwan. Plasma indium (P-In), urine indium (U-In) and creatinine adjusted U-In (U-In/Creat.) were used as internal dose of indium exposure. Plasma Krebs von den Lungen-6 (KL-6) and surfactant protein D (SP-D) were used as markers of interstitial pneumonitis. Forced vital capacity (FVC), forced expiratory volume at 1st second (FEV1), and FEV1/FVC were also evaluated by spirometry.

Results After adjusted for covariates by linear regression, plasma, urinary and creatinine adjusted indium were increased in high exposure group (P-In: β=1.13, p<0.001; U-In: β=0.54, p<0.05; U-In/Creat: β=0.63, p<0.01) and low exposure group (P-In: β=0.75, p<0.05; U-In/Creat: β=0.52, p<0.05) with comparison to reference group. Plasma KL-6 was higher in high exposure group (β=0.24, p<0.05) compared to reference group, but not for surfactant protein D (SP-D). Furthermore, FVC and FEV1 were reduced in both high exposure group (FVC: β=−0.08, p<0.01; FEV1: β=−0.05, p<0.05) and low exposure group (FVC: β=−0.06, p<0.05) compared to reference group.

Conclusion Our findings indicate indium exposure was related to restrictive lung dysfunction, decreased lung function for both FEV1 and FVC test but not for FEV1/FVC ratio. Meanwhile, increased plasma KL-6 in high exposure group also supported that indium exposure results in increased risk of interstitial pneumonitis among direct indium exposure workers. Our study provided an explanation to the consequence of indium exposure- interstitial pneumonitis-restrictive lung dysfunction.

EXPOSURE TO 2,3,7,8-TETRACHLORODIBENZO-P-DIOXIN AND MORTALITY AT A TRICHLOROPHENOL PLANT IN NEW ZEALAND

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Objectives To describe how 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) exposure influenced mortality in a cohort of workers exposed more recently, and at lower levels, than other cohorts of trichlorophenol process workers.

Methods A cohort study of 1599 men and women working between January 1, 1969 and November 1, 1988 at a plant producing the herbicide 2,4,5-trichlorophenoxyacetic acid (2,4,5-T) with TCDD as a contaminant. A toxicokinetic model developed in a previous follow up was updated to estimate cumulative TCDD exposure for each individual in the study. Calculation of cause-specific standardized mortality ratios (SMRs) and 95% confidence intervals (95%CIs) compared those never and ever exposed to TCDD. Dose-response trends were assessed firstly through SMRs stratified in quartiles of cumulative TCDD exposure, and secondly with a proportional hazards model.

Results The toxicokinetic model intercept of 5.1 parts per trillion (ppt) of TCDD was consistent with background New Zealand TCDD concentrations among older members of the population. Exposed workers had non-significant increases in all cancer deaths SMR=1.08, 95% CI 0.86–1.34, deaths from soft tissue sarcoma, SMR=2.38, 95% CI: 0.63–13.26, non-Hodgkin lymphoma, SMR=1.57, 95% CI: 0.32–4.58, diabetes, SMR=1.27, 95% CI: 0.55–2.50 and ischaemic heart disease, SMR=1.21, 95% CI: 0.96–1.50. Lung cancer deaths SMR=0.95, 95% CI: 0.56–1.53, were fewer than expected. Neither the stratified SMR nor proportional hazard analysis showed a dose response relationship.

Conclusion We found neither an excess of all cancers, or any specific cancer, nor a trend with TCDD exposure. This argues against the carcinogenicity of TCDD at lower levels of exposure.

HEALTH AND PRODUCTIVITY: A FIVE YEAR STUDY (2010–14) IN A LARGE AUTOMOBILE INDUSTRY IN INDIA

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Introduction This ‘Health and productivity’ project was implemented to identify priority health problems and health related productivity loss in a large automobile industry in India. It hopes to re-emphasize that ‘OH department’ is a partner in productivity.

Objectives
1. To estimate prevalence, incidence rates, trends and risk for health problems among employees (year 2010–14)
2. To identify leading causes of hospitalization and out-patient care among employees (year 2010–14)
3. To quantify loss in productive work time attributed to health related absence in year 2014 and forecast to year 2025

Methods A five year records analysis (2010–14) was conducted in a leading automobile industry in year 2015–16. Data was pooled from clinic visit records, annual medical examination, insurance claims and leave records, systematically using employee ID. Trends and incidence rates of leading health problems among employees were determined. Total work time (man-hours) was computed and health related loss in work-time was derived. Cox-regression was used to assess risk between work department and morbidity. ARIMA method was used to forecast productivity loss till year 2025.

Results Between 2010–14, overweight and hypertension were identified as leading health risks. Respiratory, musculoskeletal and digestive disorders were leading causes for clinic visits. Infectious diseases are leading cause for hospitalization. Health related absence accounted for 1.87% of total productive time. Forecasting indicates an increase to 9.31% by year 2025.