Comparative Pulmonary Responses Caused by Exposure to Standard Cobalt and Ultrafine Cobalt

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Abstract: Comparative Pulmonary Responses Caused by Exposure to Standard Cobalt and Ultrafine Cobalt: Qunwei ZHANG, et al. Department of Environmental Health, School of Medicine, Fukui Medical University—The aim of this study was to compare the pulmonary toxicity after exposure to standard cobalt (Std-Co) and ultrafine cobalt (Uf-Co). Rats were intratracheally instilled with 1 mg of Std-Co and Uf-Co, and wet lung weight and bronchoalveolar lavage fluid (BALF) profiles were analysed 1, 3, 7, 15 and 30 d later. The effects of Std-Co and Uf-Co on indices that can be presumed to reflect epithelial injury and permeability (lactate dehydrogenase (LDH) and total protein (TP)), and release of proinflammatory cytokine (tumor necrosis factor-alpha (TNF-alpha)) were increased throughout the 30 d post-exposure period. The results showed that the effects of Std-Co and Uf-Co on these indices were significantly higher than those of control. The results also showed that the effects of Uf-Co on indices were significantly higher than those of Std-Co from 1 to 15 d after instillation. Moreover, the number of neutrophils and LDH activity in BALF in rats after exposure to Uf-Co were significantly greater than those of Std-Co-exposed rats up to 30 d after instillation. Our findings suggest that Uf-Co has a much more toxic effect on the lungs than Std-Co, but the mechanism remains to be elucidated. (J Occup Health 2000; 42: 179–184)

Key words: Ultrafine cobalt, Standard cobalt, Bronchoalveolar lavage, Lung toxicity

Cobalt is a metal that is used in many different alloys. Inhalation exposure to cobalt causes bronchial obstruction, bronchial asthma, and interstitial pneumonitis4–6. Cobalt is also one of the constituents of hard metals, and it is widely used in industry41. Occupational exposure to cobalt and its compounds has been shown to cause lung inflammation, fibrosis, emphysema, and alveolar proteinosis7, 8. Inhalation of Uf-Co also induced macrophages damage, intracellular edema of the type I alveolar epithelium, interstitial edema and proliferation of type II alveolar epithelium9.

The toxicity of particles in the lungs is not only related to the extent of exposure but also to the particle type. For example, ultrafine titanium dioxide (Uf-TiO₂) with a diameter of 20 nm can induce a pulmonary inflammatory reaction of a greater magnitude than 250 nm TiO₂ at the same mass10–14. Ultrafine nickel and Uf-TiO₂ had also more toxicity than standard nickel and standard titanium dioxide, respectively12–16.

Ultrafine cobalt (Uf-Co) with a mean diameter of 20 nm is a new category of ultrafine particles, made by a process of vapour deposition. The characteristics of Uf-Co were described in detail previously9, 17. Our previous results showed that Uf-Co can cause sustained and substantial pulmonary inflammation17, 18. To date, however, there is no report on comparative pulmonary toxicity after instillation of Std-Co and Uf-Co. Our previous results also suggested that tumor necrosis factor (TNF-alpha) may play an important role in the pathogenic effects of Uf-Co18. And TNF-alpha is a proinflammatory cytokine produced by macrophages or mononcytic phagocytes in response to various stimuli, and the production of TNF-alpha in BALF can reflect the degree of inflammation in lungs19. Therefore, the aim of present study was to compare the toxicity of Std-Co and Uf-Co by using the bronchoalveolar lavage (BAL) technique to study the cytological and biochemical changes in the bronchoalveolar lavage fluid (BAL).
Materials and Methods

Experimental animal

Wistar male rats (specific pathogen-free), weighing 180–200 g and at 7–8 wk of age were supplied by Clear Japan, INC (JCL). The rats were housed in the animal centre of Fukui Medical University, Japan. They were fed conventional laboratory diet and had free access to food and tap water.

Materials examined

1. Uf-Co powder with an average diameter of 20 nm on was purchased as ultrafine powder from INABTA and Co., Ltd. Vacuum Metallurgical Co., Ltd., Japan (Lot No. 2237). Uf-Co with a size of <30 nm constitute more than 95.0%. The size distribution of Uf-Co was detailed previously17).

2. Cobalt powder with an average diameter of 5 μm (Std-Co) was purchased as cobalt powder from Cica Reagent, Kanto Chemical Co., Japan (Cat. No. 07385-32). The powders were suspended in physiological saline to a concentration of 1 mg/ml. The suspensions were ultrasonicated for about 30 min and then sterilised at 0.5 kgf/cm² and 120°C for 15 min.

Treatment

The rats were randomly divided into fifteen groups of 5 to 6 rats. Each experimental rat was injected intratracheally with 1 mg of Std-Co or Uf-Co in suspensions in 1 ml of physiological saline. Hereafter, rats injected with physiological saline alone were called the control group. Rats in each group were killed at 1, 3, 7, 15 and 30 d after injection.

BALF

The rats were killed by an overdose injection of phenobarbital solution (50 mg/ml) into the abdominal cavity, and the lungs and trachea were removed en bloc. The wet lung weight was measured after removing the heart, mediastinal lymphnodes and adipose tissue. The lung: the body (lung index weight ratio) was calculated. BAL was accomplished with 4×10 ml saline warmed at 37°C and massage. The recovery of bronchoalveolar lavage fluid (BALF) for each rat was measured, and the BALF recovery rate was calculated. The BALF was centrifuged at 1,500-rpm (27.67 g) for 10 min, and the first tubes were used to measure the activity of the LDH and total concentration of protein and TNF-alpha. The numbers of total cells, macrophages, neutrophils and lymphocytes were evaluated for BALF pooled from each rat.

Biochemical and cytological evaluation of BALF and biochemical evaluation in supernatant

The LDH activity in the first tube of BALF was measured with an LDH C II-test kit (WAKO Pure Chemical Industries Ltd, Japan) by the lactate matrix method20). The concentration of total protein in BALF was assessed by the Lowry method21). TNF-alpha was quantified with an enzyme-linked immunosorbent assay (ELISA) kit (Biosource International). The assay utilised a standard, 96-well quantitative immunometric “sandwich” enzyme technique according to the manufacturer’s recommendation protocol. Cells were obtained by centrifugation from all the BALF, and counted in a hemacytometer by the conventional method, and cell differentiation was done by the Wright-Giemsa method.

Statistical analysis

Results were expressed as the mean and standard error. The differences among groups were tested by one-way analysis of variance (ANOVA), followed by Bonferroni multiple comparison. If a P value was less than 0.05, the difference was considered significant. All statistics were obtained with SPSS 6.1.

Results

1. Recovery of BALF

The BALF recovery rate was more than 80%, which did not differ significantly among the experimental and control groups. Data are not shown.

2. Wet lung weight

The absolute (data not shown) and relative wet lung weights of rats instilled with 1 mg Uf-Co were significantly higher than those of the controls at 1, 3, 7, 15 and 30 d after injection; the Std-Co groups had heavier lungs than the controls except at 30 d after injection (Fig. 1).

3. Cellular and biochemical constituents in BALF

Figure 2 to Fig. 6 showed the cellular and biochemical constituents in BALF at 1, 3, 7, 15 and 30 d after instillation. Std-Co and Uf-Co instillation caused an increase in total cells, and neutrophils in BALF. Instillation Std-Co and Uf-Co caused a strong increase in the total numbers of cells, neutrophils, LDH activity, total protein and TNF-alpha in BALF at 1 day after instillation, but, these indices were decreasing within the experimental period, but at 30 d after the injection of Std-Co or Uf-Co, the total numbers of cells and neutrophils in BALF were still significantly higher than those in the control groups. The increases in total cells, and neutrophils caused by Uf-Co were marked, and at all time points were significantly greater than those of Std-Co (Figs. 2 and 3). The results also showed that Std-Co and Uf-Co increased LDH activity, total protein, and TNF-alpha in BALF compared with those of the control (Fig. 4 to Fig. 6). The LDH activity and total protein in BALF also decreased during the experimental period, and the LDH activity (except at 30 d), total protein and TNF-alpha in BALF were more significantly increased in the Uf-Co group than in the Std-Co groups.
Fig. 1. Lung: body weight ratio up to 30 d after instillation of Std-Co and Uf-Co. Values are the mean ± SE for 4 to 6 rats. * indicates significantly different from the control group, P<0.05. + indicates significantly different from the Uf-Ni group, P<0.05. Shown by ANOVA with Bonferroni comparison test.

Fig. 2. Numbers of total cells in BALF from rats up to 30 d after instillation of Std-Co and Uf-Co. Values are the mean ± SE for four to six rats. Significant differences indicated as in Fig. 1.

Fig. 3. Percentage of neutrophils in BALF from rats up to 30 d after instillation of Std-Co and Uf-Co. Values are the mean ± SE for four to six rats. Significant differences indicated as in Fig. 1.

Fig. 4. LDH activity in BALF from rats up to 30 d after instillation of Std-Co and Uf-Co. Values are the mean ± SE for four to six rats. Significant differences indicated as in Fig. 1.
Discussion

We compared the BALF inflammatory profiles after instillation of Std-Co and Uf-Co. This technique has been used to assess the induction of lung injury by mineral dusts and metallic compounds. We have presented results clearly demonstrating that Uf-Co is much more toxic to the lower respiratory tract than Std-Co when they are given in the same mass dose (1 mg).

Instillation of 1 mg Uf-Co induced pulmonary inflammation, as shown by increased lung weight sustained macrophages and neutrophil influxes, and severe injury as measured by LDH and protein in BALF. These effects of Uf-Co greatly exceeded those elicited by Std-Co. The BALF profiles suggested that this Uf-Co instillation may cause a sequence of events commencing with acute damage to the epithelia, resulting in LDH release and protein accumulation resulting from increased epithelium permeability. This may allow interstitial access of particles, leading to acute interstitial and alveolar inflammation, which are persistent because of the failure to clear the unsaturated particles.

It is generally accepted that macrophages play a key role in starting the cascade of adverse reactions to toxic particles. Neutrophils normally perform an important defensive role against microbial infection, but when macrophages increase in number and become activated in response to particle deposition, they can release chemotactic factors that attract polymorphonuclear leukocytes and monocytes. These activated macrophages and neutrophils can release a variety of cytokines, toxic oxygen metabolites and proteinase that can damage the lung parenchyma and also stimulate fibroblast proliferation.

TNF-alpha is a proinflammatory cytokine produced by macrophages or monocytes in response to various stimuli. Neutrophils are activated by TNF-alpha to a state where they are more likely to promote inflammation and cause injury to epithelial cells. Our results showed that instillation of Std-Co and Uf-Co caused a TNF-alpha increase in BALF that was greater with Uf-Co and that was persistent. And TNF-alpha in the Uf-Co group was significantly higher than that of Std-Co at all time points. Interestingly, an increase in TNF-alpha in BALF was accompanied by an increase in the total numbers of cells and neutrophils in BALF. It is suggested that TNF-alpha released from pulmonary leukocytes probably plays a role in the high toxicity of Uf-Co.

In the case of ultrafine particles, the massive number of particles may stimulate macrophages to release these mediators in exaggerated amounts. Furthermore, the total numbers of ultrafine particles may overwhelm the capacity of the macrophages to phagocytize and allow interaction of particles with epithelial cells leading to epithelia injury. Instillation with Uf-Co caused mild pulmonary damage, which was reflected by increased LDH activity and total protein in BALF. Uf-Co caused severe pulmonary damage at an early stage, but this effect...
decreased during the experiment, and was reflected by the LDH and total protein in BALF at 30 days after instillation. Nevertheless, our previous study showed that ultrafine nickel caused epithelial injury from a relatively low exposure, sufficient enough to cause death of the rats due to acute pulmonary edema\(^\text{[5]}\). Although nickel and cobalt belong to the same elemental group, ultrafine nickel was more highly and persistently toxic to lung than UF-Co\(^\text{[15]}\).

This study did not elucidate the mechanism why UF-Co is able to cause higher pulmonary toxicity than Std-Co, but the difference between UF-Co and Std-Co in particle diameter are the most striking characteristic, and UF-Co has a much greater surface area than Std-Co. Particle size appears to be very important in terms of inducing lung disease after deposition in the lungs\(^\text{[10–12, 25]}\). Furthermore, our previous studies also found that some ultrafine particles have free radical activity\(^\text{[13]}\), and exposure to UF-Co can stimulate pulmonary leukocytes to release TNF-alpha, which may underline the effects reported here.

In conclusion, our study confirmed that UF-Co is much more toxic to the lungs than Std-Co. The data obtained from this study have potential benefit for understanding the special toxicity of UF-Co. Since the present occupational exposure limit is based on mass only and does not take particle size into account, these findings have important implications for hygiene regulations.

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References

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