Hard metal lung disease: importance of cobalt in coolants

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ABSTRACT Four patients were found to react to occupational exposure to grinding of hard metal (tungsten carbide). Three of the patients had symptoms and signs compatible with an allergic alveolitis, the symptoms disappearing and the chest radiograph clearing when they were absent from work for a few months. Re-exposure to the offending agent led to new signs and symptoms. The first patient was re-exposed twice and each time reacted a little more seriously. After the last episode her chest radiograph has not cleared completely, in contrast to the first two times. The fourth patient had more typical occupational asthma. All the cases occurred in the part of the factory where air concentrations of cobalt were the lowest. The cobalt there is dissolved in the coolant necessary for grinding the hard metal. It occurs mainly in the ionised form, which is known to react with proteins and therefore presumably acts as a hapten. Protective measures, including choosing a coolant with minimal ability to dissolve cobalt and an effective exhaust system, should minimise the risk of this occupational disease in the future.

Hard metal lung disease has been described since 1940 as a cause of death in respiratory failure because of fibrosis of the lungs. Hard metal is an alloy consisting of mainly (tungsten) wolfram carbide (70–95%) and cobalt (5–25%). Various amounts of other metals, such as titanium, tantalum and so on are often added. Because of its hardness, hard metal plays an important and increasing role in the industrialised world. It was first produced in 1926 and after world war 2 production increased substantially. Sweden is, through Sandvik Coromant, one of the world's leading producers of hard metal.

Tungsten carbide has been shown to be inert, and cobalt is considered to be the cause of the damage.1–3 A considerable effort has been made over the years in the hard metal industry to reduce the levels of dust and particularly cobalt exposure in the workers' environment. The occurrence of pulmonary disease which could be connected with hard metal exposure in four workers in the present factory was, therefore, unsuspected and alarming. In this report we describe the four cases and discuss various peculiar factors in connection with them. The amount of various metals in the mediastinal lymph nodes from these patients and from control subjects is also reported.

The industrial process

The first step in the production is forming wolfram carbide (WC) from the ore. The WC is then milled to a fine powder and mixed with finely milled metallic cobalt. The purity of the powder is very important, and the size of the granules is ideally 1–2 mm. Next, the mixed powder is pressed into the roughly desired shapes and heated under pressure to about 1000°C. This gives the material enough hardness to be ground more finely for holes to be drilled into it. This is done by dry grinding. Thereafter the product is heated to about 1500°C, which is above the melting point for cobalt. Finally, the pieces are ground to very exact shapes. Since the material is very hard, only diamond grinders can be used, and cooling is necessary. Pure water cannot be used for various reasons, and different emulsions in water therefore have to be used. The compositions of various coolants are industrial secrets. The final product
consists of small pieces of metal which are very hard and durable and are used mainly in various tools for the cutting edge.

Since 1969, the Swedish TLV for cobalt has been 0.1 mg/m³. This value was often exceeded in the early 1970s, but since then this has only happened rarely. The dustiest part is the mixing area, followed by the dry grinding area where the values generally are lower. In the wet grinding area the values have been consistently very low—from unmeasurable quantities to at most 0.01 mg/m³. As a rule, the values have been from 0.002 to 0.004.

Separate investigations of the coolants by the Swedish Board for the Safety of Workers, and the Local Safety Engineer of the Factory, show that there is a tendency for the coolants to dissolve and accumulate cobalt, the amount varying a little between different compositions of the coolant. The coolant forms a fine spray of droplets which can be inhaled or deposited on the workers' skin or both. The cobalt in the coolant is mainly in the dissolved or ionised form, and after a few weeks the concentrations of cobalt had increased in some coolants up to 200 μg/g. The main difference between the "dry" and the "wet" is thus that the cobalt level is higher in the dry area but it is non-ionised, while in the wet area, the level is very low and the cobalt is in the ionised form.

Case reports

Four patients, two men and two women, were discovered to have pulmonary disease which could be connected with hard metal exposure during 1968–74. All four were working in the wet grinding area of Sandvikens Jernverk, Gimo. The period of exposure varied between two and four years (fig 1). The investigations were carried out at the Department of Pulmonary Diseases of the University Hospital, Uppsala, with technical help from the Department of Clinical Physiology and Clinical Chemistry.

Case 1

This previously healthy woman, born in 1916, had never smoked. There was no history of allergy in the patient or her relatives. In 1966, she started working in the factory in the wet grinding area. In 1968, she developed a dry cough and dyspnoea, particularly on exercise. On a chest radiograph bilateral very small infiltrates were noted. On admission to the hospital she had an intense dry cough, but no crackles could be heard over the lungs. To exclude sarcoidosis, a mediastinoscopy was performed. No treatment was given. After a few months away from work the patient felt completely healthy and the chest film was also normal. She returned to work. After a while her symptoms returned and once again she had to stop working and was reinvestigated in hospital. This time, bilateral crackles were audible and the minimal basal infiltrates had returned on the chest film. After another few months of sick leave she recovered completely again and returned to work, only to have recurrence of the signs and symptoms within a month. This time, she had a short course of corticosteroid treatment. She has not returned to work since then. Her symptoms have slowly disappeared and her chest films returned to normal after several years. Her lung function tests showed variations that closely followed her symptoms and radiographic changes (fig 2).

Case 2

This previously healthy man, born in 1904, had never smoked and had no history of atopy. He started working in the factory in 1961 in the dry area, and from 1966 in the wet grinding area (fig 1). Because of an increasing dry cough and breathlessness a chest radiograph was taken in 1969, showing bilateral small infiltrates. On auscultation, fine crackles could be heard. A bronchoscopy and a mediastinoscopy were performed to exclude sarcoidosis. All investigations revealed normal findings except the chest radiograph and the lung function tests. The patient felt better and after a few months returned to work where he promptly got worse again. This time he was treated with
corticosteroids for a short while, and slowly improved. He did not return to work but retired on an old age pension. He is still regularly checked at the hospital. At the latest visit he felt well with only very slight symptoms, but the chest film still showed minimal changes and crackles could still be heard in the lungs.

CASE 3
This previously healthy woman, born in 1916, had never smoked. There was no allergy known in the family. She started working in the factory in the grinding area in 1966, but had no symptoms until she came down with “flu” in late 1969 and after that suffered from continuous dry cough and dyspnoea. She was on sick leave from late summer 1970, and was first seen in the hospital in September 1970 when she was already much better. Bilateral crackles were heard and the chest film showed bilateral mottling. She was investigated as the other patients and since she gradually recovered, no treatment was given. After more than half a year of sick leave she returned to the factory, initially to a dry part where she suffered no symptoms, and then to the wet grinding area again. Very soon her symptoms returned and she had to be treated with corticosteroids. After recovery she has been working in a part of the factory where she is not exposed to hard metal dust. She has since remained symptomless, but her radiograph has remained abnormal with small bilateral densities without any tendency to progression.

CASE 4
This man, born in 1912, stopped smoking in 1955. There was no family history of atopy and he had been previously healthy apart from slightly raised blood pressure, treated with diazides from 1972. In 1971 he started to work in the wet grinding area of the factory and began to have asthmatic attacks on weekdays, starting Monday afternoon and evening, with remissions during weekends and holidays. The symptoms occurred only during a period of two years when he was working at a machine which lacked proper exhaust facilities. He was investigated at a time when these symptoms had subsided. The chest radiograph showed an infiltrate in the right upper lobe which may have been caused by tuberculosis or localised fibrosis. There were also small infiltrates in the left upper lobe. There were no crackles in the lungs. Inhalation of dust from the factory caused a slow but marked reduction in peak expiratory flow rate (fig 3). Peak expiratory flow rate measures during one work day even at the time when he had no overt symptoms were markedly lower (around 350 l) throughout the day than his habitual levels at the clinic or in hospital (around 450 l). The patient was moved to an area where he was not exposed to hard metal dust and remained there until retirement practically symptom-free. At the latest check-up in the lung clinic, he was symptom-free, but still had slight bilateral changes on his chest radiographs.

Bronchoscopy was performed on all four patients and was normal in all cases. All four had also a mediastinoscopy mainly to exclude sarcoidosis because of the reticular changes on the radiographs. Glands from the mediastinum of these patients as well as from four control subjects, who worked in the same factory but did not have evidence of hard metal lung disease and two unexposed control subjects, were analysed by spectrophotometric analysis at AB Atomenergi, Studsvik, Sweden according to the method described by Brune and collaborators.6 The results of these analyses are shown in table 1. The control glands in this investigation were taken from patients who, for various reasons, mainly suspicion of carcinoma, had undergone mediastinoscopy. The lymph nodes from the patients and the control subjects were all histologically normal, showing only varying degrees of anthracosis. All of the patients were non-smokers or ex-smokers, in contrast to the control subjects who were all smokers.

All four patients had contact eczema, which developed within a few months of starting to work in the wet grinding area. Patch tests were positive to cobalt in all four patients, with eczematous lesions developing within a few days after cobalt had been applied to the skin.
Lung function studies were done in all patients. Variations can be seen, closely following the degree of exposure (fig 2, table 2) DLCO was decreased in 1968 in patient 1, but had returned to normal in 1972 when she was no longer exposed.

Discussion

Hard metal lung disease was first described in 1940. Since then several other reports have followed. Earlier the disease occurred mainly in the mixing or forming areas of the factories, since these were the areas entailing the greatest risk of exposure to metal dust. Mostly the earlier reports describe a severe progressive fibrosis of the lungs in persons occupied in this type of work.7-9 Considerable evidence suggests that cobalt is the offending component.10 11 It is now well accepted that hard metal lung disease can occur in two forms: an interstitial lung disease with all the classical signs of pulmonary fibrosis and an obstructive airways syndrome.1 3 We suggest that both these seemingly different reactions actually occur through immunological mechanisms, and that they might be two variants of the same disease.

The ionised form of cobalt reacts readily with proteins12 and can therefore presumably act as a hapten. This seems to us to be the most probable explanation for the fact that the cases of lung
dissolved in observed induce fairly by Alexandersson.1
of non-specific interstitial fibrosis workers with There of working directly but symptoms, factories in metal though air even more is as well-known skin sensitiser inducing contact eczema. All our patients had contact eczema, and they had positive patch tests to cobalt. It seems probable that the ionised cobalt, which can react with proteins, is the allergenic substance inducing disease in the lung as well as in the skin, probably acting as a hapten after having been coupled to some protein in the body. Allergic contact sensitivity can be elicited apart from in the skin also in mucous membranes, and sensitised lymph nodes seem to play an important role.10 Whether the present condition represents a true allergic alveolitis or an allergic contact sensitivity in the mucous membranes of the bronchi is a matter for speculation.

None of our patients were smokers. In fact, in a recent study by Warren,17 it was found that extrinsic allergic alveolitis seemed to be significantly correlated with non-smoking in Winnipeg as well as in most reports where smoking habits were recorded.

It seems probable that the hard metal lung disease in our cases started as an allergic alveolitis,
which can progress to severe fibrosis if the exposure is not stopped. In patient 1 the reversibility of the reduced lung function was less after each exposure. It must, therefore, be important to terminate exposure immediately in cases of hard metal lung disease occurs. Since dermatological symptoms seem to precede lung symptoms it is wise to remove from exposure all who show such symptoms and have a cutaneous patch test showing sensitivity to cobalt.

Contamination of the coolants with micro-organisms, fungi, or protozoa seems unlikely as the coolants are periodically cleaned and changed; nor were any gross signs of such contamination present. We do not believe that the symptoms were caused by a variation of "humidifier fever." With a known allergen already shown to have sensitised the patient, this seems to us to be the most plausible cause of the disease. This is further strengthened by the investigations of Coates et al., who in a tungsten carbide manufacturing plant found nine patients with a respiratory sensitisation syndrome, and where tests showed cobalt to be the offending substance. Similarly, Bruckner has described a case with recurring asthma when he worked with grinding of tungsten carbide-tipped tools and where no other agents could be found to give symptoms.

The findings in the present patients led to an increased awareness about the possible harmful effects of cobalt exposure under present conditions in the Swedish hard metal industry. An epidemiological study of workers in the hard metal industry has shown that irritation of the respiratory passages is a common complaint in exposed workers and that the symptoms are mainly obstructive, but that a chronic reduction in the pulmonary function cannot be excluded. This kind of response was more common in smokers than in non-smokers. However, workers who had been removed from exposure because of complaints were not included.

There has also been extensive testing of coolants to find one which has the least tendency to dissolve the cobalt, and promising results in these aspects seem to have been reached at the factory. Likewise, there has now been much improvement in the ventilation and every machine has an exhaust system to minimise the workers' exposure. It is hoped that these measures will prevent the appearance of further cases. However, since allergic mechanisms are involved, even very minute exposures may sensitise certain individuals. So far, since our last case was diagnosed in 1973, no new cases have occurred.

References

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