Cadmium metabolism in man

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ABSTRACT Twenty-one high frequency solderers, who had been exposed to cadmium (Cd) from a solder for periods ranging from 1 month to 18 years (median 8 months; present time-weighted average 30 nmol/m³; particle size below 1μm) had Cd levels ranging from <10 to 440 nmol/l in blood and from <0.5 to 27 μmol/mol creatinine in urine. Individual workers showed considerable variations in blood Cd levels with time, but less variation in urine levels. There was a statistically significant (p < 0.001) increase of Cd in urine with increasing exposure time. Four gas solderers, who had been intermittently exposed for 8-20 years (median 17 years) had Cd levels ranging from 45 to 150 nmol/l and urine levels of from 2 to 20 μmol/mol creatinine. There was no correlation between Cd levels in blood and urine during exposure. After exposure had ceased there was a considerable decay of blood Cd in most subjects. The half-time in 11 people ranged from 25 to 146 days (median 41 days). After the decay blood levels reached a steady state. Concentrations in urine did not decrease, or did so only very slowly. There was a significant increase of levels in urine (p < 0.001) with increasing post-decay levels in blood. There was also a significant increase (0.01 < p < 0.05) of excretion of β₂-microglobulin in urine (range 1.1-18 mg/mol creatinine, median 4.7 mg/mol creatinine) measured 11-15 months after exposure had ceased, with increasing Cd levels in urine. This may indicate an effect on renal tubular function even at kidney Cd loads corresponding to Cd levels in urine of the order of as little as 10 μmol/mol creatinine.

Cadmium (Cd) is widely used in industry and its use is increasing. It is also an environmental contaminant which seriously affects the health of the inhabitants of certain areas (Friberg et al., 1974). Exposure may lead to marked accumulation in the kidney, and eventually may cause proximal tubular damage. Toxicological data are scanty in several important aspects, including the metabolism, and also with regard to those critical Cd exposures and tissue levels which adversely affect the kidney. Such data are crucial for a reliable evaluation of risks connected with exposure, and for establishing guidelines for protective measures for exposed industrial or other populations. In this study cadmium metabolism and urinary protein excretion were investigated in workers during and after occupational exposure.

Materials and methods

Subjects studied
The groups studied consisted of 21 high frequency (Hf) solderers and 4 gas solderers (Table 1). The Hf solderers worked in a typewriter factory. For

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Sex, age, exposure to Cd, and smoking habits in high frequency (Hf) and gas solderers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variables</td>
<td>Hf solderers</td>
</tr>
<tr>
<td>Number</td>
<td>21</td>
</tr>
<tr>
<td>Men</td>
<td>16</td>
</tr>
<tr>
<td>Women</td>
<td>5</td>
</tr>
<tr>
<td>Age, mean and range (years)</td>
<td>34 (19-56)</td>
</tr>
<tr>
<td>Exposure time, median and range (months)</td>
<td>8 (1-220)*</td>
</tr>
<tr>
<td>Smokers</td>
<td>15</td>
</tr>
<tr>
<td>Non-smokers</td>
<td>6</td>
</tr>
</tbody>
</table>

*Hf solderer 5 had not been exposed to Cd for 15 years.
eight hours daily, five days a week they soldered pieces to shafts using a solder containing 18-26% Cd and 40% silver, plus zinc and copper. Each worker had a smoke extractor on his bench. All Hf solderers but one had meals at their benches, usually after a hand wash. After six months' study, a non-Cd solder was substituted for the Cd solder. One of the workers (Hf solderer 5) had not been occupationally exchange to Cd for 15 years. The median exposure time was eight months.

Three of the gas solderers were employed at the typewriter factory, and one in another metal industry. Their work rooms had ventilation of varying effectiveness. Three gas solderers had meals in the work room, usually after a hand wash. The soldering was performed at irregular intervals. The exposure stopped during the study when the Cd solder was exchanged for a non-Cd solder. The median exposure time was 17 years.

**Medical Examination**

A medical history was obtained with special reference to possible effects of Cd exposure. Three Hf solderers had a history of chronic bronchitis. One gas solderer and one Hf solderer had a history of renal calculi, in both cases before onset of Cd exposure. No other renal diseases were reported. One of the gas solderers had undergone cholecystectomy prior to onset of exposure, and still occasionally felt slight pain. No other liver disease was reported. Two gas solderers had a history of recurrent anaemia. None had hypertension.

A general medical examination was performed. The blood pressures ranged from 115/60-150/90 mmHg, but in one gas solderer was 175/100 mmHg. Levels of haemoglobin in blood and creatinine in serum were within normal limits, as were the erythrocyte sedimentation rates. In all subjects but one, serum levels of AsT (aspartate aminotransferase, GOT) and AIT (alanine aminotransferase, GPT) were within the normal range. In one subject (Hf solderer 3) markedly increased levels of SAsT (10 μkat) and SAIT (16 μkat) were unexpectedly found on day 86, and he was admitted to hospital. Levels of bilirubin and alkaline phosphatase in serum were normal. Au antigen was not detected. SAsT and SAIT returned to normal after 2 months. A mild transient hepatitis was suspected. Semiquantitative determinations of albumin and glucose in urine (Albustix and Clinistix, Ames Company) were negative in all subjects.

**Cd in Blood**

Venous blood was collected in heparinised, acid-washed tubes. Duplicate analyses were performed by AAS after acid digestion of 10 ml blood samples, addition of dithizone in an ammonia-citrate buffer, and extraction of the cadmium dithizonate with methyl isobutyl ketone. The working detection limit was 5-10 nmol/l. Blood the method error in the interval 0-180 nmol/l, calculated from duplicate analyses, was in the range 5-7 nmol/l.

Ten identical samples which were analysed within a period of eight months by different operators had levels in the range 36-76 nmol/l, an arithmetic mean of 62 nmol/l, and a coefficient of variation of 18%. The accuracy of the method was confirmed by good agreement with independent procedures (AAS and

**Particle Size**

The particle size distribution was estimated by scanning electron microscopy** (Nucleopore filters, General Electric; pore diameter 0.1 μm), at a magnification of × 10 000. As large particles may be lost from the filter, their number may be underestimated.

**Cd in Air**

Representative samples of the air inhaled were collected, on several occasions during the six months before the change to non-Cd solder, on cellulose acetate membrane filters (Millipore; pore diameter 0.8 μm) with individual samplers (MSA pumps; 2-2.5 l/min). Cadmium analysis was by atomic absorption spectrophotometry (AAS) after heating the filter in 10% nitric acid or after acid digestion of the filter, addition of dithizone in buffer solution, and extraction with methyl isobutyl ketone. The overall detection limit was at worst 2.5 nmol/filter. At a sampling volume of 0.5 m3 this corresponds to 5 nmol/m3. Acid digestion of the filter after treatment with nitric acid gave an additional recovery of less than 10%. Analysis of 22 samples collected on 0.22 μm filters in the air stream after it had passed through the 0.8 μm filters, indicated a penetration through the latter of average 9% (range 0-30). Air levels reported below have been corrected for this error.

**Table 2** Cadmium levels in blood obtained by atomic absorption spectrophotometry (AAS) and by a neutron activation method (NAA)

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>AAS* (nmol/l)</th>
<th>NAA† (nmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>19</td>
<td>14</td>
</tr>
<tr>
<td>2</td>
<td>46</td>
<td>53</td>
</tr>
<tr>
<td>3</td>
<td>49</td>
<td>50</td>
</tr>
<tr>
<td>4</td>
<td>75</td>
<td>55</td>
</tr>
<tr>
<td>5</td>
<td>82</td>
<td>98</td>
</tr>
<tr>
<td>6</td>
<td>88</td>
<td>52</td>
</tr>
<tr>
<td>7</td>
<td>104</td>
<td>96</td>
</tr>
<tr>
<td>8</td>
<td>115</td>
<td>129</td>
</tr>
</tbody>
</table>

*Duplicate analyses.
†Single analyses performed at AB Isotopteknik, Stockholm, Sweden.
Table 3  Cadmium levels in urine obtained by the flame atomic absorption spectrophotometry (AAS) method used in the present study, and by a flameless AAS method.

<table>
<thead>
<tr>
<th>Sample No.</th>
<th>Flame AAS* (nmol/l)</th>
<th>Flameless AAS† (nmol/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>2</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>3</td>
<td>29</td>
<td>25</td>
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<tr>
<td>4</td>
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<td>5</td>
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</tr>
<tr>
<td>6</td>
<td>69</td>
<td>71</td>
</tr>
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<td>7</td>
<td>72</td>
<td>79</td>
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<td>8</td>
<td>85</td>
<td>84</td>
</tr>
<tr>
<td>9</td>
<td>120</td>
<td>120</td>
</tr>
<tr>
<td>10</td>
<td>203</td>
<td>204</td>
</tr>
</tbody>
</table>

*Duplicate analyses. †5 analyses in each sample.

neutron activation analysis) in an interlaboratory check (Table 2).

**CD IN URINE**

PVC bottles which had been rinsed with acid and dithizone in a buffer were used to collect 24-hour urine samples. The analytical method for Cd was the same as described above for blood, and duplicate analyses were performed. The practical detection limit was about 4 nmol/l (corresponding to about 0.3 μmol/mol creatinine). Urine analyses were in excellent agreement with results obtained by a flameless AAS method (Table 3).

**AAS APPARATUS**

All Cd determinations were carried out on a Perkin-Elmer Model 403 atomic absorption spectrophotometer using a three-slot burner head and standard operating conditions (Perkin-Elmer Corporation, 1968). A hollow cathode lamp and an electrodeless discharge lamp were used as a light source.

The method control of Cd in urine was performed on a Perkin-Elmer Model 305 B atomic absorption spectrophotometer with graphite cell and HGA 74 power supply.

**CREATININ IN URINE**

A spectrophotometric method (Teger-Nilsson, 1961) was used.* Repeated analyses of 10 identical samples gave a coefficient of variation of 2%.

**TOTAL PROTEIN IN URINE**

Total protein was determined according to Piscator (1962) in morning urine samples obtained 11-15 months after exposure had ended.†

*Analyses performed at Department of Clinical Chemistry, University Hospital, Lund, Sweden.
†Analyses performed by Dr O. Vesterberg, MD, National Board of Occupational Safety and Health, Stockholm, Sweden.

**β₂-MICROGLOBULIN IN URINE**

Determinations were performed 11-15 months after exposure had ended. The morning urine samples that had been used for total protein determination were used for β₂-microglobulin analysis. In the evening, before the urine was sampled, the worker swallowed 5 g sodium bicarbonate. The pH values of the urine samples were subsequently checked to ensure that they exceeded 5.5. A radioimmunoassay (RIA) procedure (Phadebas β₂-micro Test) was used for duplicate analyses (Pharmacia Diagnostics AB, 1975).‡ The detection limit was about 30 μg/l (roughly 2 mg/mol creatinine). The method error was 13 μg/l.

**HALF-TIMES OF CD IN BLOOD**

The kinetics of Cd in blood after exposure had ended were calculated by estimation of the constants A, B and k in the model Y(t) = A + B·e⁻ᵏᵗ using a computerised iterative procedure based on the least squares method.

**STATISTICS**

Linear and multiple regression analysis were used.

‡Analyses performed at Pharmacia Diagnostics AB, Uppsala, Sweden.

**Table 4  Cadmium concentration in air at the site of high frequency soldering**

<table>
<thead>
<tr>
<th>Cd in air (nmol/m²)</th>
<th>Exposure time* (h)</th>
<th>Percentage of total exposure time</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;10</td>
<td>186-5</td>
<td>59-0</td>
</tr>
<tr>
<td>10-19</td>
<td>49-2</td>
<td>16-0</td>
</tr>
<tr>
<td>20-29</td>
<td>36-0</td>
<td>11-0</td>
</tr>
<tr>
<td>30-39</td>
<td>9-6</td>
<td>3-0</td>
</tr>
<tr>
<td>40-49</td>
<td>4-3</td>
<td>1-5</td>
</tr>
<tr>
<td>50-99</td>
<td>17-3</td>
<td>5-0</td>
</tr>
<tr>
<td>100-199</td>
<td>1-3</td>
<td>0-5</td>
</tr>
<tr>
<td>200-299</td>
<td>5-3</td>
<td>2-0</td>
</tr>
<tr>
<td>300-399</td>
<td></td>
<td></td>
</tr>
<tr>
<td>400-499</td>
<td></td>
<td></td>
</tr>
<tr>
<td>500-599</td>
<td></td>
<td></td>
</tr>
<tr>
<td>600-699</td>
<td>7-0</td>
<td>2-0</td>
</tr>
<tr>
<td>Total</td>
<td>316-5</td>
<td>100-0</td>
</tr>
</tbody>
</table>

*85 filters.

**Table 5  Particle size distribution in smoke collected at high frequency soldering**

<table>
<thead>
<tr>
<th>Particle size (μm)</th>
<th>Percentage of particles*</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;0-1</td>
<td>62-5</td>
</tr>
<tr>
<td>0-1-1-0</td>
<td>37-3</td>
</tr>
<tr>
<td>1-1-2-0</td>
<td>0-2</td>
</tr>
<tr>
<td>&gt;2-0</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>100-0</td>
</tr>
</tbody>
</table>

*500 particles.
Results

CD IN AIR
The air concentration at Hf soldering with Cd solder seldom exceeded 90 nmol/m³ (10 μg/m³; Table 4). The time-weighted average level was about 30 nmol/m³ (3 μg/m³). Most particles were smaller than 0.1 μm, almost all smaller than 1 μm (Table 5).

The irregular frequency of gas soldering in the industries made reliable estimations of time-weighted levels of Cd in air impossible.

CD IN BLOOD AND URINE DURING EXPOSURE
Blood and urine samples were collected repeatedly during exposure. There was a considerable interindividual variation of Cd levels in blood (Fig. 1). In Hf solderers concentrations ranged from <10-440 nmol/l (<1-49 μg/l) on ten occasions over a six-month period. Considerable individual variation with time was also noted; the maximum range found was 150-440 nmol/l (17-49 μg/l) in Hf solderer 3. In urine samples obtained on five occasions levels ranged from <0.5 to 27 μmol Cd/mol creatinine (=μg/g creatinine; Fig. 1). In general the levels of Cd in urine varied less than those in blood, although in some subjects single values deviated from the individual overall concentration (for example, in Hf solderer 4). In the Hf solderers regression analysis revealed a statistically significant increase of Cd in urine when exposure time increased (n = 21; t = 7.0; p < 0.001; Fig. 2). However, the variation was considerable. In the gas solderers levels in blood ranged from 45 to 150 nmol/l (5-17 μg/l) and in urine from two to 20 μmol/mol creatinine (=μg/g creatinine; Fig. 1).

Fig. 3 shows the relationship between Cd levels
Cadmium metabolism in man

**Fig. 2** Relationship between levels of cadmium in urine and exposure time in 21 high frequency solderers (Hf S; circles) and 4 gas solderers (triangles). Closed symbols denote smokers, open non-smokers. The regression line \( y = 0.04x - 0.4 \) is calculated for Hf solderers only. Hf S 5 had not been occupationally exposed to Cd for 15 years before sampling.

**Fig. 3** Relationship between cadmium levels in blood and urine in 18 high frequency solderers (dots) and 4 gas solderers (triangles) during cadmium exposure.

in blood and urine in one day of sampling. There was no significant increase of Cd in urine with increasing levels of Cd in blood \( n = 22; t = 1.3; p > 0.05 \).

**CD IN BLOOD AND URINE AFTER THE END OF EXPOSURE**

In most workers there was a decay of blood Cd levels after the end of exposure (Fig. 1). Sufficient data to permit a closer analysis were present in 18 Hf solderers and all four gas solderers. Seven Hf solderers after a decay period reached a level of <10 nmol/l (<1 µg/l; for example Hf solderer 3). Twelve workers (11 Hf solderers and one gas solderer) reached a plateau level of 10 nmol/l (1 µg/l) or higher (for example, Hf solderers 1, 2, and 4, and gas solderer 2). In three gas solderers there was little or no decay (for example, gas solderer 1). In 11 Hf solderers in whom it was possible to analyse the decay kinetics more closely, the decay curves fitted single exponential functions (Fig. 4). The half-times (Fig. 5) in 10 workers ranged from 25 to 66 days (median 41 days), but in one worker (Hf solderer 2) was 146 days.

Changes of Cd levels in urine after end of exposure were considerably less marked than in blood. It was not possible to estimate half-times. To get a rough picture, a rectilinear regression curve was calculated in each subject on Cd in urine with regard to time after end of exposure. In 17 subjects (Hf solderers) with overall Cd levels in urine
during and after exposure below 5 μmol/mol creatinine (=μg/g creatinine), the slopes ranged from a decrease of 0.03 to an increase of 0.08 (median was an increase of 0.03) μmol/mol creatinine ·100 days⁻¹. There was a tendency (not statistically proved) to decay when levels of Cd in urine were high. In six subjects (four gas solderers and two Hf solderers) with levels of 6-16 (median 13) μmol/mol creatinine, the slopes ranged from an increase in one subject of 0.03 to a decrease of 2 (median was a decrease of 0.5) μmol/mol creatinine·100 days⁻¹. However, it must be realised that these figures are uncertain to some degree.

Fifteen months after the end of exposure there was a statistically significant (n = 25; t = 14.4; p < 0.001) increase of Cd in urine with increasing levels of Cd in blood (Fig. 6).

Proteins in Urine
In samples obtained 11-15 months after the end of exposure total protein excretion in urine ranged from 1.1-1.1 (mean 4.5) g/mol creatinine (range 10-100, mean 40 mg/g creatinine). There was no statistically significant increase in total protein excretion with increasing Cd levels in urine (n = 25; t = 1.7; p > 0.05).

The relationship between β₂-microglobulin excretion in urine (range 1.1-1.1, mean 6.7, median 4.7 mg/mol creatinine; range 10-160, mean 59, median 42 μg/g creatinine) on the one hand and Cd levels in urine (Fig. 7) and age on the other was studied by multiple regression analysis. There was a statistically significant increase of β₂-microglobulin with increasing Cd levels in urine when possible influence of age had been eliminated (n = 25; t = 2.6; 0.01 < p < 0.05), while no corresponding effect of age could be established (t = 1.1; p > 0.05). It should be noted that one subject (Hf solderer 5) had not been exposed to cadmium for 16 years before sampling. His levels of Cd in urine decreased at an approximate rate of 0.03 μmol/mol creatinine ·100 days⁻¹.

Discussion

The smoke collected from Hf soldering with Cd solder and analysed for particle size showed a predominance of very small particles. Even if the content of Cd in particles of different size is not known, it is reasonable to assume that a large part of the Cd inhaled was retained in the lungs of the workers and was absorbed into their bodies. Most of the Cd levels in air recorded in this study were low when compared to the threshold limit values (TLV) for Cd fume in workroom air in different countries (USA 50μg/m³ (440nmol/m³), (American Conference of Governmental Industrial Hygienists, 1976); Sweden 20 μg/m³ (180 nmol/m³), (Swedish National Board of Occupational Safety and Health, 1974)). As pointed out in the Materials and methods section, a minor fraction of the smallest particles may have escaped through the filters, but even taking that possibility into account, the exposures were low. The way in which Hf soldering was carried out seemed to be constant or at least fairly constant in each worker and also similar in different workers.
In spite of this, it transpired that considerable variations occurred during the air sampling periods. Fairly extensive sampling was carried out on three occasions. It is possible that even larger variations may have occurred. The reason for the variations is not fully understood, although it is clear that the size of the pieces soldered and the number of soldering operations on each shaft are important factors. It is also reasonable to assume that the condition of the ventilation equipment at each bench as well as the individual working pattern may have affected the levels, although the conditions in this respect could not be adequately mapped.

A striking feature in several Hf solderers was a decrease in blood Cd levels even before the end of exposure. The decrease started at about the time when the first results were shown to the workers, which might offer an explanation in terms of better working hygiene. However, one of the three air sampling periods occurred after the blood Cd levels had started to decrease and the air levels recorded then did not deviate compared to the earlier measurements. It is probable that a decrease of exposure occurred as a result of better hygiene with regard to eating and smoking at the work benches.

It is extremely difficult to estimate present and especially previous exposures. The working conditions in the typewriter factory had been improved by providing better work rooms and more efficient ventilation. This is of particular importance in the case of a cumulative agent like Cd. It is almost impossible to assess other routes of exposure, such as food and cigarettes contaminated by dirty hands. It thus seems advisable to rely upon analysis of biological material for the assessment of risks of poisoning.

The Cd levels in blood and urine found in this study during exposure are in accordance with findings in other industries (Friberg et al., 1974; Kjellström, 1977). Most measurements exceeded the amounts reported in non-exposed subjects by different authors (<10 nmol/l in blood and <20 nmol/day in urine).

There was an increase of Cd levels in urine with increasing duration of exposure. This might indicate a continuous accumulation but may also reflect a heavier exposure years ago.

The variations in blood Cd levels are striking: these may be due to variations in recent exposure, an assumption supported by the decrease in most subjects after exposure had ended. The levels in urine did not show the same pattern, and only a few workers had a significant but slow decrease after exposure. It is thus not surprising that there was no significant correlation between levels in blood and urine during exposure.

In many subjects it was possible to fit the decay of Cd in blood to a single exponential function. The half-time was fairly rapid and similar in different subjects but one worker deviated (Hf solderer 2). Scrutiny of his curve (Fig. 1) reveals that the levels started to decrease even before the end of exposure, probably due to reduction of exposure. If the full curve is considered, the levels decayed in a way consistent with the pattern found after the end of exposure in the other workers. It is thus probable that the long half-time obtained in that worker is due to minor analytical errors in some of the data making up the last part of the curve, but there is of course the possibility of an inter-individual variation in Cd metabolism, as has been reported in the case of methylmercury (Skerfving, 1974). Hf solderer 3, who had the highest blood Cd, had biochemical indications of mild transient liver damage (see Materials and methods). His half-time did not deviate from that of the other workers. Although liver damage has been reported in some Cd workers (Friberg et al., 1974) it seems more probable that this worker suffered from hepatitis of other origin.

The half-times of 1-2 months found in this study are considerably shorter than those reported by Piscator (data quoted by Friberg et al., 1974) in a group of heavily exposed subjects. However, a close examination of the latter curves indicates that there is no obvious disagreement if the levelling off of blood Cd seen in some workers is taken into consideration. Recently, an average half-time of 80 days was computed from data obtained in newly-employed Cd workers (Kjellström, 1977).

The rapid decay of blood Cd might reflect redistribution of one or perhaps several pools of easily movable Cd in the body. It is well known that soon after exposure a large part of the Cd is present in the liver (Friberg et al., 1974). The rapid decay might mainly reflect that pool. In a few subjects no definite shift in blood Cd occurred, possibly because of low recent exposure, and after the first rapid decay the curves levelled off. In many subjects the level obtained was consistent with concentrations in non-exposed subjects. In others the plateau reached was considerably above normal levels. These levels may reflect one or several other pools, possibly present in the kidney which is known to contain a major part of the Cd in the body some time after exposure (Friberg et al., 1974). The low background exposure also contributes to a minor extent. As it is reasonable to assume that Cd levels in urine mainly reflect concentrations in the kidneys, one would expect a good correlation between levels in blood and urine after the end of exposure, which was also the case.

Certainly a single exponential function is too
simple a model to describe the blood Cd curve. The level obtained after the initial decay is probably not the final one, as the elimination of Cd in urine in many workers considerably exceeded their assumed background exposure. It is known that the elimination of Cd from the body, mainly through the kidneys, is extremely slow; the half-time has been estimated at decades (Task Group on Metal Accumulation, 1973). It is thus not surprising that no decrease in blood Cd could be demonstrated during the limited observation period in the present workers after the first rapid decay, and that a decrease of Cd in urine was present only in a few workers and then only at a slow rate. This is in agreement with observations (Piscator, data quoted by Friberg et al., 1974) of a correlation between Cd levels in blood on the one hand and kidney damage (tubular proteinuria) on the other as much as 20 years after the end of heavy exposure.

Both total protein and $\beta_2$-microglobulin excretion in the present workers were within normal limits (Evrin and Wibell, 1972; Pharmacia Diagnostics AB, 1975). There was however a statistically significant increase in $\beta_2$-microglobulin excretion with rising Cd levels in urine, which was not explained by age. Increased elimination of $\beta_2$-microglobulin in urine is an early sign of renal tubular damage, which is in turn the first toxic effect of Cd exposure (Subcommittee on the Toxicology of Metals, 1976). Although only a small sample of workers was studied, and there is a possibility that the workers showing the highest urinary Cd levels might have had even higher Cd concentrations in urine at an earlier stage as a result of poorer working conditions, the present results may indicate a slight effect on tubular function from Cd exposure at levels in urine even of the order of 10 $\mu$mol/mol creatinine.

Data on $\beta_2$-microglobulin reported by Piscator (in Friberg et al., 1974) and Kjellström (1977) also indicate effects at low urinary Cd concentrations. Higher Cd levels have been found by several authors in workers with proteinuria (Friberg et al., 1974; Lauwerys et al., 1974; Tsuchiya, 1975). However, the methods used were less sensitive as indices of tubular dysfunction than the radioimmunological $\beta_2$-microglobulin analyses employed in the present study.

This study was supported by grant 73/134 from the Swedish Work Environment Fund. We wish to express our thanks to the workers and management of the factories studied. Statistical advice was given by Jan Rise, PhD, and Bengt Rignér, PhD. Valuable developmental work on Cd analyses was undertaken by Andreas Schütz, BSc, and technical assistance by Mrs Lisbeth Frank.

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