Effect of Rise Distance on Droplets Generated from Bubble Bursting on the Surface of Chromic Acid Solutions

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Effect of Rise Distance on Droplets Generated from Bubble Bursting on the Surface of Chromic Acid Solutions

In the processes of chromium electroplating, bursting bubbles are the major sources of chromium droplets that may lead to nasal septum disorders among workers. The gas bubbles form as a result of electrochemical reactions on the surfaces of pieces being plated, rise through the liquid, and then burst into droplets at liquid surfaces. The dimensions of the pieces being plated differ from piece to piece, and therefore the rise distance of the bubbles varies. This study is aimed at evaluating the effect of rise distance on the characteristics of droplets generated from bubbles bursting on the surface of chromic acid solutions by an experimental bubbling system. Two rise distances, 15 and 30 cm, were tested. An aerodynamic particle sizer was used to determine the droplet size distribution over the range of 0.8–30 μm. A Marple personal impactor was employed to collect droplet samples on polyvinyl chloride filters for Cr(VI) analysis. It was found that the Cr(VI) concentrations in droplets increased when the bubble rise distance increased. The result suggests that soluble ions such as Cr(VI) are scavenged by rising bubbles.

Keywords: bubbles, chromium droplets, rise distance

Control of chromium emissions is an important issue in both industrial hygiene and environmental protection. Chromium electroplating plants are the major sources of atmospheric chromium. In most chromium electroplating processes, the chromium of hexavalent form instead of trivalent form is used. Hexavalent chromium in the form of chromic acid, known as chromic acid, is more toxic than trivalent chromium.(1–6)

Epidemiological studies indicate that droplets containing Cr(VI) are probably the main cause for nasal septum disorders among chromium electroplating workers. Gomes(6) studied 303 chromium electroplating workers and found 63.4% to have nasal ulceration or perforations. Lindberg and Hedenstierna(7) examined 100 chromium electroplating workers and 119 unexposed controls with reference to changes in the nasal septum. They found that more than two-thirds of the subjects had developed nasal septum ulceration and perforation.

In Taiwan, chromium droplets generated from electroplating processes have received considerable attention because of high prevalence rates of nasal septum ulcer or perforation among workers. Lin et al.(8) conducted a survey of 7 chromium electroplating factories and examined 79 workers with reference to nasal septum lesions. They found 16 cases of nasal septum perforation and 42 with ulceration. In a separate study, Liu et al.(9) found 5 with nasal septum perforation among 45 hard chrome-electroplating workers.

In the chromium electroplating process, chromium droplets are generated from bubbles bursting at liquid surfaces. Generation of droplets from bursting bubbles is of great importance in the liquid-to-air transfer of nonvolatile compounds, which may have adverse effects on the environment and human health. Previous studies on the transfer of trace metals from seawater to air showed that oceanic droplets were enriched with trace metals. Van Grieken et al.(10) performed bubbling experiments by using radioactive tracers of metals such as 65Zn and 75Se. Their
study confirmed that the composition of resulting droplets differed from that of the bulk solution. Piotrowicz et al.\cite{11} found that trace metals in water could be highly concentrated in droplets generated through bursting of bubbles and suggested that factors such as scavenging by rising bubbles, adsorption in the surface layer, and biological processes may relate to the enrichment of these metals in droplets. However, these previous studies dealt with only dilute solutions. Little is known regarding the transfer of ions from concentrated solutions to air by the bursting of bubbles, which occurs in many industrial operations.

Data on the characteristics of Cr(VI)-containing droplets generated from bursting bubbles are needed in the control of chromium emissions and the assessment of health risk. In a practical operation the rise distance of bubbles varies because of the diverse dimensions of the pieces being plated and could have some influence on the droplet characteristics. The objective of this study, therefore, was to evaluate the effect of rise distance on the characteristics of Cr(VI)-containing droplets by an experimental bubbling system. The characteristics of droplets studied included the number size distribution, the volume distribution, and the Cr(VI) concentration in droplets.

**MATERIALS AND METHOD**

Two concentrations of chromic acid solution, 31.25 and 125 g/L, were used in this study. The change in Cr(VI) concentration in the bulk solution measured before and after each run was within 5% (in a range of 0.5 to 4.8%) even though water evaporated from the liquid surface during the bubbling process.

The experimental system, shown schematically in Figure 1, consisted of a glass cylindrical tank, a water bath, a bubbling system, and the droplet sampling/measuring instruments. A glass cylinder, 50 cm in height, was used to load the test solution for experiments. The cylindrical tank, 28.6 cm in inner diameter, was sufficiently large that the wall effect on the size of generated bubbles could be neglected. The tank was kept in a water bath to maintain the liquid temperature in the tank at 45°, the average temperature of electroplating solutions in practical applications.\cite{4} The outer wall of the tank was covered with a thermal jacket and, as a result, the air temperature inside the tank was about 37°. The humidity inside the tank was near saturation during the experiments.

The bubbling system consisted of 19 circular caps, each containing 19 glass capillaries of 0.2 mm in inner diameter and 3.2 cm length. The circular caps, made of polypropylene, were 3.5 cm in diameter. The glass capillaries were inserted into the caps, which had inside surfaces lined with a silicone pad to prevent gas leakage. The plastic caps were then mounted onto an acrylic chamber and placed at the bottom of the cylindrical tank. The bubbling system was similar to those used by Wangwongwatana et al.\cite{12} and Pilacinski et al.\cite{13}

The sparging air was obtained from a laboratory air supply, and its flow rate was regulated by a mass flow controller. The air passed through a HEPA filter and a humidifier and then entered the bubbling system, where the air passed through the acrylic chamber and emerged from the capillaries. Sparging air through the CrO3 solution in the tank led to formation of droplets above the liquid surface.

An aerodynamic particle sizer (APS, model 3310, TSI, St. Paul, Minn.) was employed to measure the number concentration and size distribution of droplets. A Marple personal impactor (model 290, Graseby Andersen, Atlanta, Ga.) was used to collect size-fractionated droplet samples for analysis of Cr(VI).

The impactor had eight stages with aerodynamic cutoff diameters of 0.6; 1; 1.5; 3.5; 6; 10; 15; and 21 μm at a sampling flow rate of 2 L/min. Droplets of various size ranges were collected on PVC filters, and the Cr(VI) concentration in each sample was determined by chemical analysis.

Limited by the height of the glass tank, only two bubble rise distances, 15 and 30 cm, were studied. An acrylic column was added on the top of the tank to provide a sampling height of 35 cm, and the outer part of the column was insulated with a thermal jacket to prevent heat loss.
In each run, the chromic acid solution of a given concentration was prepared with deionized distilled water and then heated to 45°C in a storage container that was kept in a temperature-controlled water bath. Filtered and humidified air was then introduced into the bubbling system at a given flow rate. Subsequently, the electrolyte solution was gradually transferred to the bubbling tank by a peristaltic pump (model 505S, Watson Marlow, Falmouth, U.K.) until the liquid height reached 15 or 30 cm above the capillary openings. Airborne droplets were sampled by an impactor for a duration of 2 hr. The number concentration of droplets was measured by the APS three times during the bubbling period of each run.

The U.S. National Institute for Occupational Safety and Health method 7600, developed primarily for measuring chromic acid concentrations, was used to determine the Cr(VI) concentration in each droplet sample. After sampling, each PVC filter was extracted by a few milliliters of 0.5 N H2SO4 for 5 to 10 min, and then 0.5 mL of s-diphenylcarbazide solution was added to the extract solution. The total volume of the solution was brought to 5 mL with 0.5 N H2SO4. The absorbance of the color complex was measured at 540 nm by an ultraviolet-visible (UV-VIS) recording spectrophotometer with a light path of 1 cm (model UV-160A, Shimadzu Corp., Japan).

The detection limit of the spectrophotometer was 1.8 ng/mL (equivalent to 9 ng/m³ for a 0.24 m³ air sample). Recovery of Cr(VI) sampled on a filter was experimentally determined by using a spiked aqueous solution of potassium dichromate deposited on a PVC filter. The recovery at 1 and 10 μg concentration levels was 97±3% and 95±3%, respectively, within 15 hr after spiking.

**RESULTS AND DISCUSSION**

**Droplet Volume Distributions**

The droplet volume distribution was calculated from the number distribution data obtained for all experimental conditions by the APS. Figure 2 shows the number distributions of droplets produced from bursting bubbles for various experimental conditions including the two concentrations of 31.25 and 125 g/L CrO3, two gas flow rates of 4 and 8 L/min, and two bubble rise distances of 15 and 30 cm.

The APS gave the droplet size in aerodynamic diameter, which was converted into physical diameter by the following formula:

$$d_v = \frac{d_a}{\left(\frac{\rho}{\rho_0}\frac{C}{x}\right)^{1/2}}$$

where $d_v$ is the equivalent volume diameter, $d_a$ the aerodynamic diameter, $\rho$ the droplet density, $\rho_0$ the unit density (1 g/cm³), C the Cunningham slip correction, and $x$ the dynamic shape factor. The droplet density was assumed to be identical to the density of the bulk solution. Both the Cunningham slip correction factor and the dynamic shape factor are 1 for relatively large spherical droplets measured in this study.

The droplet volume concentration in a given size interval was determined by multiplying the volume of a droplet with the geometric mean diameter by the number concentration in a given size interval ($N_{d_v}$), and $N$ the number concentration of droplets in a given size range.

Figure 3 shows that the change in bubble rise distance had a negligible influence on the volume distributions of both film and jet droplets when the gas flow rate was 4 L/min, whereas the influence became significant at a higher gas flow rate of 8 L/min.

According to the results of a previous study,(14) the film droplets and jet droplets have a dividing diameter of 7 μm. The effect of bubble rise distance on the volume concentration of droplets in the following three groups for the two rise distances are plotted in Figure 4: (a) the total droplets (larger than 0.8 μm), (b) the...
FIGURE 4. Effect of bubble rise distance on the volume concentrations of droplets in three size intervals: (a) larger than 0.8 \(\mu\)m, (b) in the size range of 0.8 to 7 \(\mu\)m, and (c) larger than 7 \(\mu\)m, for different combinations of electrolyte concentrations and gas flow rates. Each bar represents the mean of three replicates and an error bar represents one standard deviation.

Theoretical & Experimental Studies

Size-Fractionated Cr(VI) Concentrations in Droplets

The Cr(VI) concentration in droplets is of great importance in the assessment of health effects. Previous studies\(^{10,11,18-21}\) showed a fractionation effect of ions in droplets generated from bursting bubbles. The Cr(VI) concentration in droplets was calculated by dividing the airborne Cr(VI) concentration by the droplet volume concentration in a given size interval. Only the Cr(VI) concentration in the droplets of 1 to 10 \(\mu\)m in aerodynamic diameter was calculated, because the APS may gradually underestimate the droplets beyond 10 \(\mu\)m and the volume distribution mode of film droplets was mainly in this size range.

Figure 6 shows the effect of bubble rise distance on Cr(VI) concentration in droplets of 1–10 \(\mu\)m in aerodynamic diameter. The Cr(VI) concentration in droplets produced at 30-cm rise distance solution was generally higher than that at 15-cm rise distance. The difference is attributable to the scavenging effect of a rising bubble, one of the major mechanisms suggested for ion enrichment (defined as the concentration ratio of other metal ion to sodium ion) in oceanic droplets generated from bursting bubbles by Van Grieken et al.,\(^{10}\) Piotrowicz et al.,\(^{11}\) and MacIntyre.\(^{20}\)

As the rise distance of bubbles in 31.25 g/L CrO\(_3\) solution increased from 15 to 30 cm, the Cr(VI) concentration in droplets generated at a gas flow rate of 8 L/min increased more than the increase at 4 L/min. However, the Cr(VI) concentration in droplets increased by a factor of about 2 when the bubble rise distance
increased from 15 to 30 cm at gas flow rates of 4 and 8 L/min in 125 g/L CrO₃ solution. One possible explanation is that the Cr(VI) ions in droplets generated at a gas flow rate of 4 L/min in 31.25 g/L CrO₃ solution were nearly saturated and therefore the ions remained at about the same concentration as the rise distance increased.

CONCLUSIONS

For the ranges of experimental parameters studied, the following conclusions can be made.

REFERENCES


