Mutagenic Activity of Airborne Particles in Center of Metropolitan Tokyo over the Past 20 Years

Osamu ENDO1), Sumio GOTO1), Daisuke NAKAJIMA2) and Hidetsuru MATSUSHITA3)

1) Azabu University
(1-17-71 Fuchinobe, Chuo, Sagamihara, Kanagawa 252-5201, Japan)
2) National Institute for Environmental Studies
(16-2 Onogawa, Tsukuba, Ibaraki 305-8506, Japan)
3) Former University of Shizuoka
(52-1 Yada, Suruga, Shizuoka, Shizuoka 422-8526, Japan)

[Received June 18, 2015; Accepted January 12, 2016]

Summary

To estimate the risks associated with long-term human exposure to potentially mutagenic compounds, it is important to measure long-term trends in the concentrations and activity of these compounds in the environment. Samples of total suspended particulates (TSP) have been collected upon a quartz fiber filter every 6 days for the past 20 years (1980–2002) from ambient air using a high-volume air sampler placed on the rooftop of the National Institute of Public Health building in Minato, Tokyo, Japan. The mutagenicity of dichloromethane extracts of TSP samples, which divided according to season, were assayed by Ames microsuspension assay using Salmonella strains TA100 and TA98, both with and without metabolic activation system (S9 mix). Mutagenic activities (rev/m²) were calculated from the dose–response curves of the samples and from those of reference airborne particles collected by the massive sampler. Mutagenic activity increased in the first decade (1980–1990) but slowly decreased in the second decade (1990–2002). Annual average mutagenicity in TA98 was mostly higher than in TA100. Mutagenic activity was higher in autumn and winter than in spring and summer.

Key words: Salmonella mutagenicity, Microsuspension assay, Airborne particle, Long-term trend

INTRODUCTION

Ambient air contains various chemical substances, some of which are carcinogens and/or mutagens. Humans continually take these chemical substances into the body by breathing, raising concerns over chronic effects such as lung cancer caused by long-term exposure to such substances. Therefore, it is very important when considering countermeasures against lung cancer to measure yearly variation of mutagenicity of airborne particulate extracts for clarifying human exposure to carcinogens/mutagens in suspended particles in ambient air. For measuring mutagenicity of environmental pollutants, Salmonella / microsome assay (Ames assay) has been widely used because it has some advantage with sensitivity, quantitativity, repeatability, and handling in comparison with other methods1). Microsuspension assay2), one of modifications of Ames assay, is more sensitive and needs less sample volume than original method. It is also easier to compare the test result with original method because it has same principle and employ same tester strains with original method. There have been therefore so many reports on mutagenicity of environmental pollutants with microsuspension assay3–19). Although there was only a previous study of mutagenicity of airborne particulates over quarter century in Sapporo, a large city in Hokkaido region northern Japan10), there was no study in Tokyo, largest city and capital of Japan.

In the present paper, the mutagenic activity of airborne particles in center of metropolitan Tokyo over the past 20 years was measured by using a sensitive method, microsuspension assay.

MATERIALS AND METHODS

Reagents

Solvents: Pesticide residue analysis grade dichloromethane (DCM; Kokusan Chemical Works Ltd., Japan) and fluorometric analysis grade dimethylsulfoxide (DMSO; Dojin Chemical Laboratory Ltd., Japan) were used.

Metabolic activation system: S9/Cofactor A set for Ames test (S9 mix; Oriental Yeast Co. Ltd., Japan) was used.

Sampling and extraction

Total suspended particulate (TSP) samples have been collected upon quartz fiber filter (20 x 25 cm, Pallflex Products Co., U.S.A.) using high volume air sampler (Kimoto Electric Co., Japan) on the rooftop of former National Institute of Public Health located in the central part of Tokyo metropolitan (Shirokanedai, Minato-ku), Japan over the past 20 years (from March 1980 to February 2002 every 6 days (occasionally 4–7 days, including very few lack) for 24 hours), at a flow rate of about 1.3 m³/min. The filter samples were stored in a
deep-freezer (−80°C) wrapped in aluminum foil and then in a plastic bag until extraction of organic matter. The filter samples were taken, and divided according to the seasons: spring (March, April and May), summer (June, July and August), autumn (September, October and November) and winter (December, January and February). TSP samples were collected and stored every 6 days so each group included approximately 15 daily samples. Two disks 40 mm in diameter were cut out from each collected filter using a belt punch, and the disks were cut into small pieces and placed into a 500 mL Erlenmeyer flask with a screw cap. Then, 150 mL of dichloromethane (Pesticide residue analysis-grade) was added to the flask and extraction was performed twice for 20 min each in a sonication bath. After sonication extraction, the supernatant fluid was filtered into an evaporation flask and concentrated in vacuo to about 150 mL with an evaporator. The volume of the condensate was measured with a 200 mL volumetric flask and 5 mL of condensate was moved to a vial for PAH analysis. The remain extraction (195 mL) was moved again to evaporation flask and also concentrated in vacuo to about 2 mL with an evaporator and the condensate was moved to a vial with a screw cap for mutagenicity test. After the solvent was removed by evaporation under a weak nitrogen gas flow, those samples were stored in a deep-freezer at −80°C until their mutation assay.

**Mutagenicity test**

The mutagenicity test was conducted by a microsuspension procedure which was a slight modification of Kado’s method2) using Salmonella typhimurium TA98 and TA100 strains (x 20 conc. bacterial solution: high-speed refrigerated centrifuge Hitachi CF15R) under the conditions of both with and without a metabolic activation system (S9 mix). The extracts were dissolved in DMSO and put into test tubes with 10, 5, 2.5, and 1.25 µL in duplicates for each dose. After adding 100 µL of 0.1 M sodium phosphate buffer or S9 mix, and 100 µL of concentrated bacteria, the tubes were capped and preincubated at 37°C with shaking. After 90 minutes of preincubation, the tubes were removed and 2.5 mL of molten top agar containing both 0.5 mM biotin and 0.5 mM histidine were added. The molten suspensions were immediately mixed with a vortex mixer and poured onto minimal glucose plates. The plates were incubated at 37°C in the dark for 48 hours and were counted using an automatic laser colony analyzer (Spiral System Instruments Inc., Model 500A). Mutagenic activity was calculated from the slope of the linear portion of the dose-response curve using the statistical model of least squares linear regression.

**RESULTS**

According to the reports of IPCS12-16), it was suggested that inter- and intra-laboratory variances were existed due to bioassay conditions, such as tester strain, metabolic activation, material, technical error and so on. These reports were also suggested that complex mixture reference materials such as NIST SRMs were very useful to reduce these variances (repeatability and reproducibility) for biological study of complex environmental samples. In our present study, it was necessary to perform the assay is divided into 13 times for all samples that were made. For the purpose to reduce the variances, an independently established reference material (RM) sample which is DMSO solution of extract from airborne particles collected by massive air sampler on the rooftop of NIPH between 1997 and 2000 has been provided17). This RM has been prepared for some research project grant awarded by Ministry of the Environment18).

Table 1 shows mutagenic activity of RM sample and correction factor for normalization of mutagenic activity of airborne particulate samples. Correction factors were calculated from the mutagenic ac-

<table>
<thead>
<tr>
<th>Test No.</th>
<th>TA100 (−S9)</th>
<th>TA100 (+S9)</th>
<th>TA98 (−S9)</th>
<th>TA98 (+S9)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>rev/µL CF</td>
<td>rev/µL CF</td>
<td>rev/µL CF</td>
<td>rev/µL CF</td>
</tr>
<tr>
<td>1</td>
<td>15.05 (1.156)</td>
<td>10.84 (0.795)</td>
<td>46.06 (1.077)</td>
<td>17.94 (1.055)</td>
</tr>
<tr>
<td>2</td>
<td>12.28 (0.934)</td>
<td>16.28 (1.194)</td>
<td>46.52 (1.088)</td>
<td>20.28 (1.193)</td>
</tr>
<tr>
<td>3</td>
<td>13.28 (1.020)</td>
<td>23.37 (1.714)</td>
<td>25.40 (0.594)</td>
<td>18.21 (1.071)</td>
</tr>
<tr>
<td>4</td>
<td>11.16 (0.857)</td>
<td>13.46 (0.987)</td>
<td>21.01 (0.491)</td>
<td>8.48 (0.499)</td>
</tr>
<tr>
<td>5</td>
<td>9.55 (0.733)</td>
<td>10.34 (0.758)</td>
<td>39.59 (0.926)</td>
<td>15.87 (0.933)</td>
</tr>
<tr>
<td>6</td>
<td>11.33 (0.870)</td>
<td>13.30 (0.976)</td>
<td>34.84 (0.815)</td>
<td>9.95 (0.585)</td>
</tr>
<tr>
<td>7</td>
<td>13.09 (1.005)</td>
<td>12.91 (0.947)</td>
<td>42.04 (0.983)</td>
<td>13.19 (0.776)</td>
</tr>
<tr>
<td>8</td>
<td>15.38 (1.181)</td>
<td>14.02 (1.028)</td>
<td>50.57 (1.183)</td>
<td>24.92 (1.465)</td>
</tr>
<tr>
<td>9</td>
<td>14.02 (1.076)</td>
<td>11.12 (0.816)</td>
<td>55.90 (1.308)</td>
<td>19.77 (1.163)</td>
</tr>
<tr>
<td>10</td>
<td>12.90 (0.990)</td>
<td>10.36 (0.760)</td>
<td>44.90 (1.050)</td>
<td>22.65 (1.332)</td>
</tr>
<tr>
<td>11</td>
<td>15.03 (1.154)</td>
<td>13.64 (1.001)</td>
<td>52.35 (1.225)</td>
<td>22.14 (1.302)</td>
</tr>
<tr>
<td>12</td>
<td>13.70 (1.052)</td>
<td>12.32 (0.904)</td>
<td>42.28 (0.989)</td>
<td>11.50 (0.676)</td>
</tr>
<tr>
<td>13</td>
<td>12.55 (0.964)</td>
<td>15.27 (1.120)</td>
<td>54.28 (1.270)</td>
<td>16.17 (0.951)</td>
</tr>
</tbody>
</table>

mean ± sd 13.02 ± 1.69 (1.000) 13.63 ± 3.45 (1.000) 42.75 ± 10.55 (1.000) 17.01 ± 5.09 (1.000)

CF (correction factor) was ratio of the mutagenic activities of each assay and their averages. For example, CF of TA100 (−S9) at the test No.1 (1.156) = 15.05 / 13.02 (mean value of 13 tests)

Negative (solvent = DMSO) control values were 226 +/- 35 for TA100 (−S9), 225 +/- 39 for TA100 (+S9), 57 +/- 21 for TA98 (−S9), and 53 +/- 16 for TA98 (+S9), respectively. A2 and B2 were used as positive controls. Mutagenic activities (revertants per micro gram) of A2 were 1300 +/- 1760 for TA100 (−S9), 2360 +/- 489 for TA98 (−S9), and those of B2 were 677 +/- 82 for TA100 (+S9), 82 +/- 29 for TA98 (+S9), respectively.
Fig. 1 Representative dose-response results for mutagenicity of airborne particulate samples by microsuspension assay

- ○ TA100 (−S9) - △ TA98 (−S9) - ● TA100 (+S9) - ▲ TA98 (+S9)

Fig. 2 Yearly variation of mutagenic activity of TSP per volume of ambient air in Tokyo from 1980 to 2001 for strains TA100 and TA98

- □ without S9mix - ■ with S9mix
been relatively low and has moderately decreased in every years.

Figure 3 shows long-term seasonal variation of mutagenic activities of TSP per volume of ambient air. In this figure, mutagenic activity was indicated in revertants per unit volume of air (revertants / m³). And also open square (□), open rhombus (◇), closed square (■), and closed rhombus (◆) indicate spring, summer, autumn, and winter respectively. From the figure, a clear seasonal variation was obtained in 1980s without regards to tester strains either with or without S9mix. Seasonal variation has become smaller after 1990. Generally, the activities in autumn-winter were higher than those in spring-summer. Moreover, the activities in autumn-winter season have dramatically decreased regardless to tester strains either with or without S9mix, although those in spring-summer season have moderately decreased for TA98 strain.

Figure 4 shows yearly variation of mutagenic activity of TSP per weight of TSP in Tokyo from 1980 to 2001. From the figure, the mutagenic activity for TA98 without S9mix has shown the maximum in 1983 and almost decreased in years after. However the activities for TA100 strain (both with and without S9mix) and those for TA98 with S9mix have shown almost even flat status. As similar as Fig. 2, the activities for TA98 strain have been higher than those for TA100 under the condition without S9 mix in every year. Under the condition with S9 mix, most of samples have obtained similar results in 1980s, although some activities for TA100 have rather been higher than those for TA98 after 1990.

Figure 5 shows long-term seasonal variation of mutagenic activi-

---

Fig. 3  Long-term seasonal variation of mutagenic activities of TSP per volume of ambient air

- □ - spring  ◇ - summer  ■ - autumn  ◆ - winter
ties of TSP per weight of TSP. In this figure, the activity was indicated in revertants per unit weight of particle (revertants/mg, particle). The legends were as same as Fig. 3. From the figure, as similar as Fig. 3, a clear seasonal variation was obtained without regards to tester strains and with/without S9/mix. However, the variation has not always decreased after 1990. The activities in autumn-winter were higher than those in spring-summer regardless to tester strains either with or without S9/mix. Overall, seasonal variation of mutagenic activity per unit weight of particle was larger than that per unit volume of air. This reason is not clear currently. These result suggested that not only TSP concentration decreased during these 20 years, but also that the generation source of the pollutant has changed.

**DISCUSSION**

Matsumoto et al. reported mutagenic activity of airborne particulates over quarter century (1974–2001) in Sapporo, largest city in Hokkaido region northern Japan[18, 29]. According to the report, mutagenicity and BaP concentrations showed seasonal fluctuations, highest in winter, lowest in summer, reflecting fuel consumption and meteorological factors. On the other hand, Tokyo has fewer differences of the temperature than Sapporo. It was suggested that these differences of the meteorological factors reflected the difference of mutagenic aspects in Tokyo and Sapporo. Long-term trends in BaP indicates dramatic reduction, while indirect-acting mutagenicity showed a moderate decline and direct-acting mutagenicity showed neither a district increase nor decrease[18-20]. Nitro-aromatic compounds have been increasing yearly in proportion to the number of diesel-powered vehicles. Those compounds play a major contributory role in mutagenic air pollution in Sapporo[19, 21]. Tokyo is the capital and has biggest populations in Japan. The aspects of mutagenicity such as yearly and/or seasonal variations of airborne particulates in Tokyo were very similar to those in Sapporo. From a comparison study, microsuspension assay showed about 7 times higher sensitivity than that of preincubation assay[21]. Our present results showed 1.34 (TA100+S9) – 5.96 (TA98 – S9) times higher sensitivity than those of Matsumoto et al. in overall mean value. It seems to be a similar tendency with more considerations when it is early for ten years. It is thought that the tendency to decrease of mutagens in airborne particulates have been successful by cumulative of many pollution-abatement measures.

We have previously reported that the concentrations of PAHs decreased in past 20 years (1980–2001), especially in the late 1980s, and slowly decreased from the middle 1990s[22]. These PAHs were also higher in autumn/winter seasons than in the spring/summer seasons. These results were very similar as the report of Matsumoto et al. and as our present results of mutagenic activity. Moreover, according to the data from Tokyo Metropolitan Environmental Agency (TMEA), the concentration of sulfur dioxide, nitrogen dioxide and suspended particulate matter in the air at Shirokanedai station (near by our sampling point) also showed seasonal variation[23].

Also according to the reports of TMEA[23], yearly average concentration of SPM had decreased from the beginning of successive measurement (fiscal year 1973) up to around 1979. Before then, although not in successive measurement, there were some difference of the concentrations of airborne particles collected with HV sampler by sampling points (industrial > commercial > residential). However it has seesawed and has been no difference by sampling points since 1980[23-24].

From our present results, it was suggested that the successive tendency to decrease of the mutagenicity of airborne particulates was caused by the effects of automobile exhaust measures and by the moves of the factories to foreign countries such as Southeast Asia areas and so on. However, it has also been suggested that take a long time (20–30 years or more) for appearance of lung cancer by mutagens/carcinogens in air. From those facts, it was suggested that more detailed studies to chase a secular variations of the qualitative changes of the automobile exhaust with the changes of the fuel in future. Authors plan to measure the various kinds of harmful compounds such like nitro-aromatics in the specimen samples of airborne particles from now on.

**ACKNOWLEDGEMENT**

This research was partially supported by a research project grant awarded by the Azabu University.

**REFERENCES**


9) Bombick, B.R., Avalos, J.T., Nelson, P.R., Conrad, F.W. and Doolittle, J.: Comparative studies of the mutagenicity of environmental tobacco smoke from cigarettes that burn or primarily heat

---

Fig. 5 Long-term seasonal variation of mutagenic activities of TSP per weight of ambient air

- □ - spring
- ◇ - summer
- ■ - autumn
- ◆ - winter


