TSP, PM10 and PM2.5 Distribution Characteristics in the Thermal Power Plants in Korea

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Abstract: In this study, the emission characteristics and heavy metal contents of TSP, PM10 and PM2.5 pollutants from three thermal power plants in Korea were investigated and compared to the electric production capacity, type of fuel and sort of air-pollution-control device. For the measurement and analysis, Korean standard test method US EPA method were used. The average concentration of TSP, PM10 and PM2.5 emitted from Plant A were 7.39, 6.16, 4.83 mg/Sm$^3$, Plant B was 5.82, 4.87, 2.35 mg/Sm$^3$ and Plant C was 1.54, 1.40, 10.02 mg/Sm$^3$, respectively. Plant A that uses heavy oil as the main fuel showed higher TSP, PM10 and PM2.5 than Plant B that uses mostly anthracite coal, and plant B showed higher TSP, PM10 and PM2.5 than Plant C that mainly uses bituminous coal. The concentration of fine particles decreased as electricity-production capacity increased. The fractions of PM10 and PM2.5 in TSP were relatively high in tested plants; this result means that more fine particles than coarse particles were emitted from all stacks. The distribution of heavy metals by particle size showed similar trends in all plants. The concentration of Zn and Mn in TSP, PM10 and PM2.5 showed higher than the others in all plants. These results confirm that the content of heavy metals in the particulate matter is influenced by the fuel that the plant uses.

Keywords: Emission, TSP, PM10, PM2.5, Fuel, Heavy Metals, Thermal Power Plant

1. Introduction

Atmospheric pollutants can directly or indirectly harm human health and properties, and ecosystems. These air pollutants are classified into Criteria Air Pollutants (CAPs) and Hazardous Air Pollutants (HAPs) [1]. These substances can be emitted by natural events such as volcanic eruptions, forest fires and outgassing from swamps, or by human technologies such as static facilities or vehicles.

Particulate pollutants are generated by mechanical treatment such as crushing and sorting of a substance, or by combustion and decomposition processes. Fine particles include secondary dusts such as fume, soot, carbon black, combustion nuclides, and oxygen-containing hydrocarbons, sulfates and nitrates. Coarse particles include solid particles such as cement dust, coal dust, and liquid particles such as raindrops, sprays, and fogs.

Management of particulate pollutants has been intensified, and attention has been focused on the effects of particle size and of the components that constitute each particle. In 1983, criteria for Total Suspended Particulates (TSPs) were introduced in Korea. In 1995, the air environment standard for the particulate matter of diameter ≤ 10 µm (Particulate Matter 10, PM10) was strengthened (to an average of 100 µg/m$^3$ for 24 h; average of 50 µg/m$^3$ for a year). From 2015, Korea has begun to manage particles with diameters ≤ 2.5 µm (Particulate Matter 2.5, PM2.5), and the management standard is based on an average of 50 µg/m$^3$ for 24 h or an average of 25 µg/m$^3$ for a year [1]. According to data collected in 2014, 91.8% of the 255 measurement sites in the nation exceeded the 24-h average and 38.8% exceeded the PM10 annual average [2]. The average annual concentration of PM2.5 in the six major cities in Korea in the last three years has been 32 µg/m$^3$ (28-39 µg/m$^3$), which exceeds the US standard (12 µg/m$^3$) [3] and the
European standard (25 µg/m³) [4], and is about three times the level of 10 µg/m³ recommended by WHO [5]. Studies on human effects on PM10 have reported an increase in mortality of about 0.5% per 10 µg/m³ of PM10 [6-7]. Based on these data, WHO recommends an annual average concentration of PM10 of 20 µg/m³, and a 24-h average concentration of 50 µg/m³ for short-term exposure. Because of its small size, PM2.5 is not filtered by the bronchi, and weakens pulmonary function or causes cardiovascular diseases [8]. Long-term exposure to PM2.5 concentration of 11–15 µg/m³ has associated risks, so WHO recommends an annual average of 10 µg/m³ [9-11].

In Korea, the necessity of managing PM2.5 emerged in the 1990s, as patients with respiratory-related diseases and those with cardiovascular-related diseases died early due to the effects of PM2.5 inhalation [12]. To avoid these consequences, the sources of PM2.5 must be identified, and plans to reduce their emissions must be developed. According to the domestic air pollutant emission data in 2014, total TSP emission amount is about 147 ton/y, PM10 about 98 ton/y and PM2.5 about 63 ton/y. Point sources of pollution emit 63 – 83% TSP, 17 – 36% PM10 and 0.2 – 0.7% PM2.5 [13].

Among point sources, manufacturing combustion processes emit the highest levels of TSP, PM10, and PM2.5. However, combustion facilities of the energy-generation industry emit TSP that have larger proportions of PM10 or PM2.5 than do combustion facilities of manufacturing processes. Therefore, production of fine dust should be managed in combustion facilities of the energy-generation industry sources. Pollutants generated during coal combustion especially particulate matter below PM10, can harm human health [14-16]. These pollutants are particularly problematic in developing countries, where dust from coal combustion accounts for one-third of the total amount of dust generated [17].

In September 2017, the Korean government announced the "Plan for Fine Dust Management" jointly with the related ministries to plan detailed implementation of measures to reduce fine dust emission from old coal-fired power plants. The highest priority is to replace or upgrade the facilities to reduce production of fine dust by coal-fired power plants. The first step toward meeting this goal is to accurately measure and characterize the particulate and particulate precursors generated by the current operating facility. Also, the components of fine dusts must be identified. Many existing studies have stated that harmful heavy metals in fine dusts can exacerbate affect respiratory diseases and cardiovascular diseases [18].

Therefore, in this study, emission characteristics and constituents of TSP, PM10 and PM2.5 were investigated in three operating thermal power plants in Korea. The plants were selected to have different electric production capacity, fuel and sort of the air pollution control devices; the distributions of fine particle and contents of heavy metals in fine particles were investigated according these differences.

2. Methods and Materials

2.1. Target Plants

Generally, a thermal power plant burns by injecting fuel with air into the boiler. A selective catalytic reduction system (SCR) or a nonselective catalytic reduction system (SNCR) is used to reduce the concentration of NOx in the emitted combustion gases. An electrostatic precipitator (ESP) or a fabric filter (FF) is used to remove dust. A flue-gas desulfurization system (FGD) is used to remove SOx.

![Figure 1. Process and condition of each plant tested.](image)

In this study, three thermal power plants with different capacity and used fuel were selected as target plants (Figure 1). Plant A has a capacity of 20 MW/h and uses heavy oil as the main fuel (33603 L/d), with 724 ton/d of anthracite coal, 1384 ton/d of bituminous coal and 1275 L/d of diesel as sub-fuels. Plant B has capacity of 200 MW/h and uses anthracite coal as the main fuel (975 ton/d), with 244 ton/d of bituminous coal and 137 L/d of heavy oil as sub-fuels. Plant C has a capacity of 500 MW/h and uses bituminous coal as the only fuel (4993 ton/d). Plant A and B used SNCRs. Plant C had a separate denitrification facility (SCR).

2.2. Sampling Method

PM10 and PM2.5 were sampled (Figure 2) in accordance with the air pollution test standard "ES 0112.1 - Method of collecting particulate matter of exhaust gas" [19] and "ES 01317.1 - Method of collecting fine dust (PM10 and PM2.5)" [20]. The test method is applicable only when the temperature of the final exhaust gas is ≤ 260°C, and should satisfy constant velocity suction coefficient of 90-110%. The PM10 and PM2.5 are removed from TSP by a cyclone combination unit, then collected using a nozzle directly in front of the filter holder. The PM10 cyclone had cut diameter (D50) = 9 – 11 µm, and the PM2.5 cyclone had D50 = 2.25 - 2.27 µm. The sampling rate was varied according to the exhaust gas temperature. In this case, a nozzle capable of collecting particles within the range of D50 should be used [21]. Sampling was performed three times in the stack that is the final outlet of each plant. The sample was collected using a circular quartz filter which was heated at 500°C for 2-3 h. The samples volumes were > 2 Sm³, but differed slightly depending on the temperature, the dynamic pressure and the static pressure of the exhaust gas of each plant.
2.3. TSP, PM10, PM2.5 Concentration Calculation and Heavy Metal Analysis Method

2.3.1. TSP, PM10, PM2.5 Concentration Calculation Method

The filter holder (figure 2) was placed in a quartz filter, samples were collected, and the filters were stored in individual filter cases for transport to the laboratory. In the laboratory, the moisture was removed and the weight concentration of particulates was determined using a balance (model: METER MT, UMT) that can measure with a precision of $10^{-4}$ g. The dust concentration in the exhaust gas was calculated by dividing the weight concentration by the sampled volume of gas.

2.3.2. TSP, PM10, PM2.5 Heavy Metal Analysis Method

The investigated target compounds were Cd, Cr, Cu, Mn, Ni, Pb, V and Zn. Most of these have emission standard values [mg/Sm$^3$]: Cd, 0.02; Cr, 0.3; Cu, 5; Ni, 2; Pb, 0.2 and Zn: 5 [22]. Mn and V in particulate matter have no emission standard yet, but they are abundant in coal and heavy oil. For the analysis of the target compounds in the collected particulate matter, EPA method 3051A [23] was used. In the pretreatment method, 10 mL of nitric acid was injected into a microwave sample pretreatment apparatus (model: MARS, CEM), and about 15 mL of distilled water was added so that the filter was immersed. The vessel was placed in a hood to remove the generated gas, then disassembled using a microwave sample pretreatment apparatus. The dissolved sample was filtered through filter paper (Whatman No. 41), then the sample solution was prepared by adding the sample to a 50-mL volumetric flask that was then filled to the mark with distilled water. The target compounds were analyzed using Inductively Coupled Atomic Emission Spectrometer (820-MS, Varian). A standard heavy-metal stock from Accustandard was used for heavy-metal analysis. The standard solution was prepared by diluting a standard stock solution of 1000 mg/L to 1, 3, 5, 10, or 20 µg/L, and adjusting the nitric acid concentration to 1%. Samples were analyzed according to EPA 200.8 9 [24]. A rinse blank was used between samples. The accuracy and precision experiments the methods were performed according to the QA/QC Handbook [25]. The accuracy was 80-90%, which is sufficient. The precision was 2.6-16.9%, which is less than the tolerable range of 20%, so the reproducibility in the experiment is confirmed. The calibration curves of target compounds all had coefficients of determination ($r^2$) > 0.9998. The linear range of the calibration curve was used to check the stability of the instrument and the conditions.

3. Results and Discussion

3.1. Concentration of TSP, PM10 and PM2.5

The concentration and fraction of TSP, PM10 and PM2.5 were calculated for each plant (Table 1). Generally, the TSP is calibrated to 6% oxygen for power plants that use solid fuels or liquid fuels [22]. However, in this study, the TSP is calibrated to 6% oxygen for power plants that use solid fuels or liquid fuels [22]. In this study, the measured values are presented without oxygen correction to facilitate comparison of PM10 and PM2.5.

Plant A had TSP = 4.62 - 9.78 (7.39 avg.) mg/Sm$^3$, PM10 = 5.02 - 8.35 (6.16 avg.) mg/Sm$^3$ and PM2.5 = 3.16 - 6.71 (4.83 avg.) mg/Sm$^3$. The fraction of PM10 in TSP was ~83%, the fraction of PM2.5 in TSP was ~65%, and the fraction of PM2.5 in PM10 was ~78%.

Plant B had TSP = 5.43 - 6.11 (5.82 avg.) mg/Sm$^3$, PM10 = 4.45 - 5.40 (4.87 avg.) mg/Sm$^3$ and PM2.5 = 2.03 - 2.80 (2.35 avg.) mg/Sm$^3$. The fraction of PM10 in TSP was ~84%, the fraction of PM2.5 in TSP was ~40%, and the fraction of PM2.5 in PM10 was ~48%.
Plant C had TSP = 1.46 - 1.60 (1.54 avg.) mg/Sm$^3$, PM10 = 1.31 - 1.57 (1.40 avg.) mg/Sm$^3$ and PM2.5 = 0.96 - 1.13 (1.02 avg.) mg/Sm$^3$. The fraction of PM10 in TSP was ~90%, the fraction of PM2.5 in TSP was ~66%, and the fraction of PM2.5 in PM10 was ~73%.

The distribution of fine dust varies according to the emission plant, but also varies depending on the capacity and process of the plant and the fuel used [18]: in combustion plants that used fossil fuels, TSP was in the range of 0.7-14.4 mg/Sm$^3$, and PM10 was 88.4-97.0%, PM2.5 was 65.7-75.5% and PM10 was 25.5-52.0%; when the fuel was anthracite coal, the fraction of PM10 and PM2.5 to TSP was somewhat low, and at all of the plants, the dust collector was equipped with an electrostatic precipitation [18]. In the present study, the highest concentration of TSP was found in Plant A that uses heavy oil. Of TSP in this plant, 83.4% was PM10 and 65.3% was PM2.5. In addition, TSP concentration was higher in Plant B that uses anthracite than in Plant C that uses bituminous coal. Of TSP in Plant B, 83.6% was PM10 and 40.4% was PM2.5. TSP concentration in emissions was lowest from Plant C, but in it 90.2% was PM10 and 66.1% was PM2.5. This is the same trend as noted by Ehrlich et al. [18] and the same as in surveys of the fine dust emitted by domestic thermal power plants [26-29]. In addition, Ehrlich et al. [18] reported that plants that have an SNCR that inject elements for NOx treatment have higher TSP concentrations than plants that lack an SNCR. Our results agree with these.

In this study, TSP concentrations were higher for Plants A and B, which used an SNCR to remove NOx, than for Plant C, which uses an SCR. The amount of particulate matter was highest in the plant that had the smallest. Therefore, this study demonstrated that the distribution of particle concentration and particle size are affected by the capacity and process of the plant, the type of prevention plants, and the fuel used, as suggested by Ehrlich et al. [18]. The distributions of emitted TSP, PM10 and PM2.5 also differed among the plants (figure 3). The distributions of PM10 and PM2.5 were normalized by setting TSP to 100%; ~90% of TSP was PM10 and ~66% of TSP was PM2.5 in all three plants.

### Table 1. Mean and (range) of concentration [mg/Sm$^3$] and fraction [%] of TSP, PM10 and PM2.5 in samples from each plant.

| Plant | TSP     | PM10    | PM2.5    | PM10/TSP | PM2.5/TSP | PM2.5/PM10 |
|-------|---------|---------|----------|----------|-----------|------------|
| A     | 7.39 (4.62-9.78) | 6.16 (5.02-8.35) | 4.83 (3.16-6.71) | 83.4 (65.8-108.6) | 65.3 (47.1-86.4) | 78.3 (55.2-131.2) |
| B     | 5.82 (5.43-6.11) | 4.87 (4.45-5.40) | 2.35 (2.03-2.80) | 83.6 (77.8-91.1) | 40.4 (34.2-51.6) | 48.3 (37.5-62.9) |
| C     | 1.54 (1.46-1.60) | 1.40 (1.31-1.57) | 1.02 (0.96-1.13) | 90.2 (83.0-98.2) | 66.1 (60.2-71.6) | 72.7 (61.1-86.2) |

[Figure 3. Distribution of TSP, PM10 and PM2.5 emitted from the stack of the tested plants.]

### 3.2. Concentration of Heavy Metal in TSP, PM10 and PM2.5

#### 3.2.1. TSP

The concentrations of heavy metal in TSP differed among the plants (Table 2, Figure 4). The total concentration of heavy metals in TSPs decreased in the order Plant A > Plant B > Plant C. The compositions were as follows: Plant A had Zn > Mn > Cr > Pb > V > Ni > Cu > Cd; Plant B had Mn > Zn > Ni > V > Cu > Pb > Cr and Plant C had Mn > Zn > V > Cu > Ni > Pb > Cd > Cr. Mn, V, and Zn are mainly emitted in solid fuel combustion plants, and Cr, Ni, and Cu are mainly emitted in liquid fuel combustion plants [30-31]. Plant C uses only bituminous coal as fuel; its results are similar to those of Pio et al. [30-31] and Querol et al. [32]. Plant B showed a different...
tendency, possibly because it uses solid and liquid auxiliary fuels. The higher Mn and Zn in Plants B and C than in Plant A were attributed to the use of coal as the main fuel; the higher Pb, Cr, and Ni in Plant A than in Plants B may occur because Plant A uses heavy oil. In addition, the higher Mn and Zn in Plant A than in Plants B and C may be the effects of its auxiliary fuels, anthracite and bituminous coal. The difference in the content of heavy metals in the three plants seems to be due to the difference in fuel used.

### Table 2. Mean and (range) of concentrations [µg/Sm³] of heavy metal in TSP, PM10 and PM2.5 in each plant.

| Element | Plant | Size | Cd  | Cr  | Cu  | Mn  | Ni  | Pb  | V   | Zn  |
|---------|-------|------|-----|-----|-----|-----|-----|-----|-----|-----|
|         | A     | TSP  | 0.06 | 1.05 | 0.77 | 3.13 | 0.85 | 1.01 | 0.95 | 3.25 |
|         |       | PM10 | 0.03 | 1.69 | 0.59 | 3.33 | 1.55 | 0.88 | 0.95 | 3.19 |
|         |       | PM2.5| 0.08 | 1.99 | 0.81 | 2.53 | 1.07 | 0.69 | 0.76 | 2.78 |
|         | B     | TSP  | 0.04 | 0.44 | 0.47 | 6.29 | 0.59 | 0.46 | 0.50 | 3.83 |
|         |       | PM10 | 0.03 | 0.25 | 0.43 | 6.06 | 0.34 | 0.44 | 0.50 | 3.56 |
|         |       | PM2.5| 0.03 | 0.16 | 0.30 | 5.63 | 0.45 | 0.30 | 0.42 | 3.27 |
|         | C     | TSP  | 0.01 | 0.18 | 0.20 | 3.08 | 0.14 | 0.12 | 0.27 | 0.71 |
|         |       | PM10 | 0.00 | 0.00 | 0.15 | 3.38 | 0.20 | 0.12 | 0.27 | 0.41 |
|         |       | PM2.5| 0.01 | 0.20 | 0.22 | 2.64 | 0.06 | 0.09 | 0.20 | 0.77 |

3.2.2. **PM10**

The concentrations of heavy metal in PM10 differed among the plants (Table 2, Figure 5). The total concentration of heavy metals in PM10 for each plant was in the order Plant A > Plant B > Plant C, as with TSP. In Plant A the order was Mn > Zn > Cr > Ni > V > Pb > Cu > Cd; in Plant B it was Mn > Zn > V > Pb > Cu > Ni > Cr > Cd, and in Plant C it was Mn > Zn > V > Cu > Ni > Pb > Cd > Cr. PM10 showed high concentrations of Mn, Zn and V, and low concentration of Cd, as with TSP. The rankings for each of these compounds were somewhat different, possibly as a result of the difference in fuel used, as is the case of TSP. The distribution tendency of heavy metals in PM10 for each plant is similar to the distribution of heavy metals in TSP for each plant. However, in PM10 from Plant A, the concentration of Ni was higher than that of V, and in PM10 of Plant B the concentration of Ni was slightly lower than that of V, compared to TSP. That the difference in fuel affects the distribution of heavy metal, and these heavy metal can also be distributed differently depending on the particle size.
3.2.3. PM2.5

The concentrations of heavy metals in PM2.5 also differed among the plants (Table 2, Figure 6). The total concentration of heavy metals in PM2.5 for each plant was in the order Plant A > Plant B > Plant C, as with TSP and PM10. Plant A the following was: Zn > Mn > Cr > Ni > Cu > V > Pb > Cd; in Plant B it was Mn > Zn > V > Pb > Cu > Ni > Cr > Cd, and in Plant C it was Mn > Zn > V > Cu > Pb > Ni > Cd > Cr.

The rankings for each of these compounds also differed, possibly as a result of differences in the fuel used, as in the cases of TSP and PM10. Although the tendency of distribution of heavy metal in PM2.5 is similar to that in TSP and PM10, Plant A showed the highest concentration of Pb, and slightly higher Zn than Mn compared to TSP and PM10. The heavy metals seem to show different distribution tendencies by particle size.

4. Conclusion

This study quantified the distributions of TSP, PM10 and PM2.5 and contents of heavy metals in those particles in emissions from three thermal power plants, which use the different fuel types and have the different electric-production capacities. The following conclusions were obtained.

(1) The emission concentration of TSP, PM10 and PM2.5 were affected by the type of fuel used and the electric generation capacity. These measurements were all higher in emissions from Plant A that uses heavy oil as the main fuel, than in emissions from Plant B that uses mainly anthracite, or in emissions from Plant C which uses mainly bituminous coal as fuel; the
measurements were also higher in Plant B than in Plant C. The concentrations of particulate matter decreased as the electricity-generation capacity of the plant increased.

(2) The percentage of PM10 and PM2.5 to TSP differed among plants. The fraction of each particle size relative to the total concentration of particulate matter is the PM10 for TSP in the plant using bituminous coal. The fraction of PM2.5 was high. In the plants that use bituminous coal, the emissions of TSP were low, but had high proportions of PM10 and PM2.5. However, in the other plants, the emission of TSP was high had high proportions of PM10 and PM2.5. The efficiency of the electrostatic precipitator installed in each plant may be too low for removal of fine particles.

(3) The distribution of heavy metal by particle size of each plant was similar, but the distribution of heavy metals obviously differed among the plants. All of three measures of particulate matter were influenced by the fuel used. Therefore, the distribution characteristics of heavy metal according to particle size are different according to the type and amount of fuel used.

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