Investigation of Aerosol Number Concentration at Gosan Site in Jeju, Korea

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(Manuscript received 12 December, 2011; revised 20 December, 2011; accepted 25 January, 2012)

Abstract

The aerosol number concentration have measured with an aerodynamic particle sizer spectrometer (APS) at Gosan site in Jeju, Korea, from March 2010 to March 2011. And then the atmospheric aerosol number concentration, the temporal variation and the size distribution of aerosol number concentration have been investigated. The aerosol number concentration varies significantly from 748 particles/㎝³ to zero particles/㎝³. The average number concentration in small size ranges are very higher than those in large size ranges. The number concentrations in the size range 0.25–0.28 ㎛, 0.40–0.45 ㎛ and 2.0–2.5 ㎛ are about 84 particles/㎝³, 2 particles/㎝³ and 0.4 particles/㎝³, respectively. The number concentrations in range of larger than 7.5 ㎛ are below 0.001 particles/㎝³.

The seasonal variations in the number concentration for smaller particle (<1.0 ㎛) are not much, but the variations for larger particle are very evident. And strong amplitudes of diurnal variations of entire averaged aerosol number concentration are not observed.

Size-fractioned aerosol number concentrations are dramatically decreased with increased particle size. The size-fractioned aerosol number concentrations in size range 0.8–4.0 ㎛ during nighttime are evidently higher than during daytime, but similar levels are appeared in other size range. The seasonal differences in the size-fractioned number concentrations for smaller size range (<0.7 ㎛) are not observed, however, the remarkable seasonal differences are observed for larger size than 0.7 ㎛.

Key Words: Aerosol, Number concentration, Seasonal variation, Diurnal variation Size-fractioned concentration, Gosan site

1. Introduction

Atmospheric aerosols are emitted from a large variety of sources, thus they have various sizes, shapes and made up of hundreds of different chemicals. Aerosols have an important influence on the atmospheric visibility, irradiation balance and they also act as cloud condensation nuclei and affect precipitation patterns. Therefore atmospheric aerosols play an important role in the variation of weather and climate. There is growing concern related to the human health effects of atmospheric aerosols, because of the epidemiological studies that the adverse health effects of atmospheric aerosols include respiratory irritation and changes in pulmonary function. (Vyziene and Girgzdys, 2009; Sharma et al., 2003)

The size distribution, composition, sources and sinks of atmospheric aerosols are important parameters in understanding and managing aerosol effects on health, visibility and climate. The size distribution and composition of atmospheric aerosols, especially,
are key factors in assessing these effects. Recently, there has been an increased interest in the relative health effects of particles of smaller sizes. Several researchers have also reported that the health effects of aerosols, for a given mass concentration, are larger for smaller particles. (Buzorius et al., 1999; Stanier et al., 2004; Bigi and Ghermandi, 2011) The number concentration of small size aerosol is usually very high, but its mass concentration is relatively low. Several researches have also shown that the health effects of aerosols may be more sensitive to the number than mass concentration. (Yan et al., 2004) In spite of these facts, most of the air quality managements so far have been based on aerosol mass concentration.

Recently, a large number of observational studies on aerosol number concentrations have been performed. The features and temporal and/or spatial variations of atmospheric aerosol number concentrations have been investigated. (Vyziene and Girgzdys, 2009; Sharma et al., 2003; Stanier et al., 2004; Buzorius et al., 1999; Choi et al., 2005) Many studies on the relationship of aerosol number concentration and mass concentration have been performed. (Yan et al., 2004; Bigi and Ghermandi, 2011) A lot of studies on the aerosol number concentrations during Asian dust storm have been done. (Watanabe et al., 2005; Zhang et al., 2008; Chun et al., 1999) And the meteorological dependences of aerosol number concentration have been analyzed. (Paatero et al., 2005; Hussein et al., 2006) Also the number concentrations of atmospheric ultrafine aerosol have been studied. (Sharma et al., 2011; Longly et al., 2005; Minoura and Takekawa, 2005) However, few continuous measurements have been made of size-separated number concentration of atmospheric aerosol in Jeju, Korea.

In this study, we have continuously measured the aerosol number concentration with an aerodynamic particle sizer spectrometer (APS) measuring from 0.25 to 32.0 μm from March 2010 to March 2011 at Gosan in Jeju. The temporal variation of size-separated number concentration of atmospheric aerosol and the size distribution of aerosol number concentration have been studied.

2. Materials and Methods

2.1. Monitoring site

The monitoring site, Gosan (33°17’N, 126°10’E), is located at the western tip of Jeju Island. The site, shown in Fig. 1, is one of the background sites in Korean Peninsula. Jeju Island is a volcanic island with peak elevation of ~2000 m and is a major resort area with no large industrial sources. The island is located in the East China Sea, ~100 km south of the Korea mainland, ~250 km west of Kyshu, Japan, ~500 km east-northeast of Shanghai, China. The site is about 70 m above sea and adjacent to the seashore.

![Fig. 1. Location of the monitoring site and surrounding region.](image)

2.2. Instruments and measurements

In the present research, an aerodynamic particle sizer spectrometer (APS, GRIMM Aerosol Technik
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Fig. 2. The grand average aerosol number concentration measured separately in the 30 size ranges.

GmbH & Co., Model #179), which uses light scattering method, has been installed at Gosan site. This instrument measures atmospheric aerosol number concentration in 30-channel of different size range from 0.25 ~ 0.28 μm to 30.0 ~ 32.0 μm.

Atmospheric aerosols have been sampled at 3 m from the ground level and the number concentration have been measured after having removed moisture. The measurement of the aerosol number concentration has been performed every 5 minute, from March 2010 to March 2011. The number concentration of aerosol particles have been measured separately in the 30 size ranges, such as 11-channel in range of 0.25 ~ 1.0 μm, 4-channel in range of 1.0 ~ 2.5 μm, 8-channel in range of 2.5 ~ 10.0 μm, 7-channel in range of 10.0 ~ 32.0 μm. The 1-hour averaged data of the number concentration of aerosol obtained every 5-minute from APS have been used in this study.

3. Results and Discussions

3.1. Overview of aerosol number concentration

The grand average aerosol number concentrations measured separately in the 30 size ranges during the experimental period are shown in Fig. 2. From these results, it is found that the aerosol number concentration varies significantly with particle size from 748 particles/cm³ to zero particles/cm³. The average number concentrations in small size ranges are higher greatly than those in large size ranges. The average concentration is about 84 particles/cm³ in the smallest size range (0.25 ~ 0.28 μm), 15 particles/cm³ in the size range 0.40 ~ 0.45 μm, 2 particles/cm³ in the size range 0.58 ~ 0.65 μm, 0.4 particles/cm³ in the size range 2.0 ~ 2.5 μm. On the other hand, the average number concentrations of larger particles (≥7.5 μm) are lower than 0.001 particles/cm³.

The values of overall aerosol number concentrations have large differences according to measurement methods, in particular, lower limit measuring size range. In general, the values of overall aerosol number concentrations are greatly more than 10,000 particles/cm³ in case of using the method which can measure ultrafine particles (≤100 nm). (Sharma et al., 2011; Longly et al., 2005; Minoura and Takekawa, 2005) However, in this study, the overall aerosol number concentration in excess of 1000 particles/cm³ has never observed because the lower limit measuring size range (0.25 ~ 0.28 μm) is large, relatively.
The aerosol number concentrations measured in this study are low in comparison with other previous researches performed in urban area. For example, Sharma et al. has reported that the aerosol number concentrations measured in Roorkee, India, in the size range 0.3-0.5 μm, 1.0-2.0 μm and 2.0-5.0 μm are $1.5 \times 10^4$-$2 \times 10^3$ particles/ℓ, $1 \times 10^3$-$5.6 \times 10^3$ particles/ℓ and $50$-$300$ particles/ℓ, respectively. (Sharma et al., 2003) Also, Park et al. has reported that the aerosol number concentrations measured in Seoul, in the size range 0.3-0.5 μm, 0.82-2.23 μm, 2.23-6.06 μm and 6.06-10.0 μm are $32.3$ particles/cm³, $8.5$ particles/cm³, $1.6$ particles/cm³ and $0.008$ particles/cm³, respectively. (Park et al., 2001)

Table 1. shows the basic statistics of the number concentration data in the five wide size ranges (particle/cm³)

<table>
<thead>
<tr>
<th>Size Range(μm)</th>
<th>Mean</th>
<th>Min</th>
<th>Max</th>
<th>S.D</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25–0.5</td>
<td>241.488</td>
<td>3.149</td>
<td>1703.365</td>
<td>234.085</td>
</tr>
<tr>
<td>0.5–1.0</td>
<td>8.413</td>
<td>0.033</td>
<td>189.152</td>
<td>11.867</td>
</tr>
<tr>
<td>1.0–2.5</td>
<td>1.543</td>
<td>0.008</td>
<td>260.466</td>
<td>9.484</td>
</tr>
<tr>
<td>2.5–5.0</td>
<td>0.810</td>
<td>0.003</td>
<td>249.195</td>
<td>8.514</td>
</tr>
<tr>
<td>5.0–10.0</td>
<td>0.014</td>
<td>0.000</td>
<td>9.325</td>
<td>0.148</td>
</tr>
</tbody>
</table>

3.2. Temporal variation of aerosol number concentration

The number concentration of ambient particles changes not only with the season but also within minutes. In this study, the seasonal averages of the aerosol number concentration for six size ranges are shown in table 2 to find the seasonal variation of those.

Table 2. The seasonal averages of the aerosol number concentration for six size ranges (particle/cm³)

<table>
<thead>
<tr>
<th>Size Range (μm)</th>
<th>Spring</th>
<th>Summer</th>
<th>Fall</th>
<th>Winter</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25–0.5</td>
<td>251.974</td>
<td>215.690</td>
<td>232.388</td>
<td>254.578</td>
<td>241.488</td>
</tr>
<tr>
<td>0.5–1.0</td>
<td>9.288</td>
<td>8.722</td>
<td>7.251</td>
<td>8.540</td>
<td>8.413</td>
</tr>
<tr>
<td>1.0–2.5</td>
<td>3.165</td>
<td>0.842</td>
<td>1.042</td>
<td>0.834</td>
<td>1.543</td>
</tr>
<tr>
<td>2.5–5.0</td>
<td>2.197</td>
<td>0.208</td>
<td>0.363</td>
<td>0.228</td>
<td>0.810</td>
</tr>
<tr>
<td>5.0–10.0</td>
<td>0.036</td>
<td>0.004</td>
<td>0.009</td>
<td>0.004</td>
<td>0.014</td>
</tr>
<tr>
<td>10.0–32.0</td>
<td>0.002</td>
<td>0.000</td>
<td>0.001</td>
<td>0.000</td>
<td>0.001</td>
</tr>
</tbody>
</table>
The seasonal averages of the aerosol number concentration in size range of 0.25–0.5 μm are from 215.7 to 254.6 particles/cm³, those in size range of 0.5–1.0 μm are from 7.3 to 9.3 particles/cm³ approximately, hence the seasonal differences are less than 24% of overall averages. However, the average number concentrations in size ranges of larger than 1.0 μm during the spring time are much more than other season. For example, the spring time average in size range of 2.5–5.0 μm is more than ten times the summer time average. It can be seen from table 2 that the seasonal differences in the number concentration for smaller particle ranges(0.25–0.5 μm, 0.5–1.0 μm) are not much, but the differences for larger particle ranges are very evident. It is well established that airborne particles over size 1 μm are dominated by mineral particles with quite different sources and transport properties than finer particles(〈1 μm). (Longly et al., 2005) It seems that the seasonal average number concentrations for larger particles in spring are significantly high due to Asian dust storm, pollen etc., on the other hand those in summer are low due to scavenging by precipitation. Stanier et al. have reported that there is the absence of a clear seasonal trend in average aerosol number concentration in small size range(〈0.56 μm). (Stanier et al., 2004)

To find diurnal variation of the aerosol number concentration, the hourly averages of the concentration during experimental period are presented in Fig. 3. The diurnal aerosol number concentration pattern is driven by emission characteristics of dominant sources and meteorological conditions. The aerosol number concentration has a evident diurnal variation in urban
area due to the local sources of finer aerosol. In general, the aerosol number concentration is high during rush hour and nighttime. (Yan et al., 2004; Vyziene and Girgzdys, 2009; Sharma et al., 2003; Bigi and Ghermandi, 2011) However, it is can be seen from Fig. 3 that strong amplitudes of diurnal variations of entire averaged aerosol number concentration are not observed because of the location of measurement site with no large anthropogenic sources.

As shown in Fig. 3, in the size ranges 0.25 ~ 0.5 μm and 0.5 ~ 1.0 μm, the aerosol number concentration during daytime is slightly lower than nighttime. This trend in spring season is appeared remarkably compared to other season. It is suggested that the fine aerosol is generated by condensation because of high humidity and low temperature at that time. (Sharma et al., 2003) The similar diurnal variation patterns to these results are reported previously by many other researchers. (Yan et al., 2004; Vyziene and Girgzdys, 2009; Bigi and Ghermandi, 2011) On the other hand, interesting diurnal variations of the aerosol number concentration are found in the size ranges 1.0 ~ 2.5 μm and 2.5 ~ 5.0 μm. The aerosol number concentrations during daytime (especially, 10:00 ~ 16:00) in the spring season are drastically lower than nighttime, though there are no evident diurnal variations of the concentrations in the summer, fall and winter. The reason of this phenomenon is not clearly realized in this study, thus the intensive studies are required to explain this tendency in future.

3.3. Size distribution of aerosol number concentration

It is known that the aerosol composition and size distribution are key factors in assessing the effects on human health, global heat balance and weather. (Bigi and Ghermandi, 2011)

Size-fractioned aerosol number concentration, \[ N(D) = dN/d\log(D) \], for the entire averages, for the daytime averages and the nighttime averages and for the seasonal averages are presented in Fig. 4., Fig. 5., and Fig. 6., respectively. As shown in Fig. 4., it is found that the \( N(D) \) values for the entire averages are dramatically decreased with increased particle size, although three different patterns are observed with the size range. For example, the \( N(D) \) curve for smaller size range(<0.8 μm) has a steep slope but the \( N(D) \) curve for size range 0.8 ~ 3.0 μm has a more gentle slope, and then the \( N(D) \) curve is changed steeply for larger size range(>3.0 μm).

Fig. 5. shows that the differences in size distributions for smaller size range(<0.8 μm) and for larger size range(>4.0 μm) between the daytime averages and the nighttime averages are not much. However, in size range 0.8 ~ 4.0 μm, the \( N(D) \) values during daytime are evidently higher than during nighttime. It is assumed that the aerosol group in size range 0.8 ~ 4.0 μm has a slight number variation but has an evident temporal variation with size range.
The seasonal differences in the size-fractioned number concentrations for smaller size range (<0.7 μm) are not observed, however, the seasonal averages of the N(D) values for larger than 0.7 μm are remarkably different (Fig. 6). For example, the size-fractioned number concentrations during the spring are significantly higher than other season, and the N(D) values in coarse particle ranges (>10 μm) during the summer are greatly low compared to other season. It is seemed that these tendencies caused by the sources and sinks of larger particles, such as, Asian dust storm and pollen in spring and scavenging by precipitation in summer.

4. Conclusions

The aerosol number concentration have measured with an aerodynamic particle sizer spectrometer (APS) at Gosan site in Jeju, Korea, from March 2010 to March 2011. And then the atmospheric aerosol number concentration, the temporal variation and the size distribution of aerosol number concentration have been studied. Results derived from this study are summarized as follows.

The aerosol number concentration varies significantly from 748 particles/cm³ to zero particles/cm³. The average number concentration in small size ranges are very higher than those in large size ranges, such as, about 84 particles/cm³ in the smallest size range (0.25 ~ 0.28 μm), 0.4 particles/cm³ in the size range 2.0 ~ 2.5 μm and below 0.001 particles/cm³ in range of larger than 7.5 μm.

The seasonal variations in the number concentration for smaller particle (<1.0 μm) are not much, but the variations for larger particle are very evident. And strong amplitudes of diurnal variations of entire averaged aerosol number concentration are not observed, but interesting diurnal variations are observed in the size ranges 1.0 ~ 2.5 μm and 2.5 ~ 5.0 μm. The concentrations during daytime are drastically lower than nighttime in the spring, but there are no evident diurnal variations in other season.

Size-fractioned aerosol number concentrations are dramatically decreased with increased particle size, although the curve slope varies with the size range. In size range 0.8 ~ 4.0 μm, the size-fractioned aerosol number concentrations during nighttime are evidently higher than during daytime, but similar levels are appeared in other size range. The seasonal differences in the size-fractioned number concentrations for smaller size range (<0.7 μm) are not observed, however, the remarkable seasonal differences are observed for larger size than 0.7 μm.

Acknowledgements

This work was funded by the Korea Meteorological Administration Research and Development Program under Grant RACS 2011-1015.

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