Beryllium-7 Aerosols in Ambient Air

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Abstract

The aerodynamic size distribution of $^7$Be aerosol particles in ambient air was determined by using 1-ACFM and high-volume (HVI) cascade impactors, 20 m above the ground on the roof of the Faculty of Science building, Aristotle University of Thessaloniki at Thessaloniki Greece (40°38’N, 22°58’E) from November 2006 to June 2008. The activity concentration of $^7$Be aerosols was determined by gamma spectrometry ($E_\gamma = 0.477$ MeV). The activity size distribution of $^7$Be-aerosols was largely associated with submicron aerosol particles in the accumulation mode (0.4-2.0 μm). The activity median aerodynamic diameter, AMAD varied from 0.76-1.18 μm (average 0.90 μm), indicating post-condensation growth either in the upper atmosphere or after mixing into the boundary layer. The geometric standard deviation, $\sigma_g$ varied from 1.86-2.77 (average 2.24). The activity size distribution of $^7$Be-aerosols peaked in the 0.7-1.1 μm size range in the 65% of the measurements carried out by the 1-ACFM cascade impactor. In estimating lifetimes of $^7$Be-aerosols in ambient air, a mean residence time of about 8 days averaged for atmospheric aerosols of 0.90 μm AMAD size.

Keywords: Beryllium-7; Radioactive aerosols; Aerosol impactors; 1-ACFM; AMAD; Residence time.

INTRODUCTION

Beryllium-7 is a relatively short-lived ($T_{1/2} = 53.3$ days, $\tau = 1/\lambda = 77$ days) naturally occurring radionuclide of cosmogenic origin which is formed by spallation processes of light atmospheric nuclei, such as carbon ($Z = 6$), nitrogen ($Z = 7$) and oxygen ($Z = 8$), when the above nuclei absorb protons and even neutrons of the primary component of cosmic rays, according to the following reactions:

\[
\begin{align*}
^{12}\text{C}^6_p &\rightarrow ^8\text{Be}^3\text{Li} \\
^{14}\text{Ne}^3_p &\rightarrow ^7\text{Be}^2\text{He} + ^2\text{He} \\
^{28}\text{Si}^p &\rightarrow ^7\text{Be}^3\text{He} + ^3\text{He} \\
^{12}\text{C}^7_n &\rightarrow ^8\text{Be}^3\text{He} \\
^{14}\text{N}^7_n &\rightarrow ^7\text{Be}^3\text{Li} \\
^{28}\text{Si}^7_n &\rightarrow ^7\text{Be}^3\text{Li} + ^3\text{He} \\
\end{align*}
\]

Essentially, the $^7$Be atoms are produced by high-energy spallation interactions. Production
is greatest in the upper stratosphere, but some energetic cosmic-ray neutrons and protons survive in the lower atmosphere, producing cosmogenetic radionuclides, such as $^7$Be. Production is not only altitude- but also latitude-dependent and varies as well with the 11-year solar cycle that modulates cosmic-ray penetration through the earth’s magnetic field. The calculated global average production rate of cosmogenic $^7$Be per unit surface area of the earth is 810 atoms/m$^2$/s and the average concentration of $^7$Be in the troposphere is 12.5 mBq/m$^3$ (UNSCEAR 2000).

Once $^7$Be is formed in the troposphere, it rapidly associates primarily with submicron-sized aerosol particles (Bondietti et al., 1984; 1987). Beryllium-7 in these fine aerosols may subsequently enter the marine as well as the terrestrial environment and vegetation via wet or dry depositional events. Following deposition, $^7$Be will tend to associate with particulate material (particle-reactive element).

Beryllium-7 has come to be recognized as a potential tool in studying the description of environmental processes, such as aerosol transit and residence times in the troposphere (Martell 1970), aerosol deposition velocities (Young and Silker 1980) and aerosol trapping by above ground vegetation (Bondietti et al., 1984).

Beryllium-7, and other natural radionuclides like $^{22}$Na, $^{32}$P, $^{33}$P, $^{35}$S and $^{210}$Pb, participate in the formation and growth of the accumulation mode aerosols (0.07-2 μm diameter) which is a major reservoir of pollutants in the atmosphere (Bondietti et al., 1987). Following its production by gas-phase nuclear transformation, this isotope condenses on the aerosol population, growing by condensation of non-radioactive species e.g. sulfates or organic (McMurry and Wilson, 1982; 1983) and so the fate of $^7$Be will become the fate of the carrier aerosols (Bondietti et al., 1984; 1987). On the behavior of $^7$Be atoms with atmospheric aerosols, it was concluded from early aerosol studies that considerable coagulation occurred during migration of $^7$Be atoms from the stratosphere and upper troposphere to ground level air (Friedlander, 2000; Grundel and Porstendörfer, 2004; Grundel et al., 2005).

This paper summarizes results of an investigation designed to characterize the aerodynamic size distributions and the aging of atmospheric aerosols in the context of $^7$Be distributions for better understanding of aerosol growth mechanisms and the behaviour of radioactive aerosols in the atmosphere.

**EXPERIMENTAL PROCEDURE**

The aerodynamic size distribution of $^7$Be in atmospheric aerosols was achieved by using Andersen 1-ACFM cascade impactors. This 1-ACFM impactor involves a flow rate of 1.7 m$^3$/h (28.3 L/min or 1 ft$^3$/min) and eight atmospheric pressure stages for collecting aerosols above 0.4 μm (Fig. 1). The effective cutoff diameters (ECDs) of this impactor were as follows: 0.4, 0.7, 1.1, 2.1, 3.3, 4.7, 5.8 and 9.0 μm. The stainless steel plates supplied by the manufacturer were used for aerosol collection. Glass fiber filters were used as back up filters to collect particles below the
Fig. 1. A schematic diagram of an 1-ACFM cascade impactor complete system.

0.4 μm collection plate. Polycarbonate films can also be used as back up filters.

The length of each collection period was 1 week. The samples were collected 20 m above the ground on the roof of the Faculty of Science building, Aristotle University of Thessaloniki, Greece (40°38’N, 22°58’E).

The stainless steel collection plates of the impactor as well as the back up filters used as plane radioactive sources were measured for ⁷Be activity (E_γ = 0.477 MeV) by gamma-ray spectrometry using a high resolution (1.9 keV at 1.33 MeV ⁶⁰Co), high efficiency (42%), low-background high purity Ge coaxial-type detector. Uncertainty in counting of ⁷Be activity was varied from 10-20%. Less than 3 mg of particulate material on any impactor stage was collected in each sampling, thus overloading (≥ 10 mg on any stage) was excluded. A typical gamma-ray spectrum of a glass fiber air filter obtained by a Ge detector, in which the 0.477 MeV γ-ray peak of ⁷Be is clearly shown in Fig. 2.

High-volume cascade impactors, HVI can also be used for the size fractionation of atmospheric aerosol particles. These impactors have a regular air-flow rate either of 0.57 m³/min (20 cfm) or 1.13 m³/min (40 cfm) and the effective cutoff diameters (ECDs) are 0.41, 0.73, 1.4, 2.1, 4.2 and 10.2 μm for the 20 cfm configuration or 0.49, 0.95, 1.5, 3.0 and 7.2 μm for the 40 cfm configuration at the standard temperature and pressure.

Fig. 2. Plot of a γ-ray spectrum of an atmospheric aerosol sample (air filter) obtained by a Ge detector.
atmospheric conditions (25°C and 760 mm Hg).

RESULTS AND DISCUSSION

Beryllium-7 Aerosol Size Distribution

A typical plot of the activity size distribution of 7Be versus aerodynamic diameter (D_p) is represented in Fig. 3. This distribution was selected from 11 measurements (samplings) made over an almost 2-y period.

Atmospheric aerosol size distribution appears to follow a trimodal distribution expected for condensation-derived aerosols. This trimodal distribution of atmospheric aerosols is showing the following modes: (i) the Aitken nuclei mode which ranges from 0.003-0.07 µm (average 0.015 µm), (ii) the accumulation mode which ranges from 0.07-2 µm (average 0.3 µm) and (iii) the coarse mode which ranges from 2-36 µm (average > 10 µm) (NRC 1979). Young et al. (1975) reported that 7Be is attached primarily to submicron-sized particles in the atmosphere. About 88% of 7Be was found to be present on particles smaller than 1.1 µm in diameter, and less than 1% was on particles larger than 7 µm in diameter. That means, 7Be-aerosols are accumulation mode aerosols. It is also evident from the plot of Fig. 3.

From eleven measurements carried out in a 2-y period at Thessaloniki, Greece with a temperate latitude (40°38’N, 22°58’E) precipitation-free (dry) climate, the activity median aerodynamic diameter (AMAD) varied from 0.76-1.18 µm (average 0.90 µm) and the geometric standard deviation (σ_g) varied from 1.86-2.77 (average 2.24). The AMAD and σ_g calculations were made by plotting the cumulative distributions on log-normal probability paper. It was shown that 69% of the 7Be activity was associated with aerosol particles with diameter smaller than 1.1 µm.

Table 1 shows data for the activity median aerodynamic diameter, AMAD of atmospheric aerosol particles associated with 7Be atoms. The AMAD values for 7Be-aerosols reported from the literature varied from 0.33-1.15 µm with the upper values being pretty close to the upper value of this work.

From a different view of the obtained results, the 7Be activity size distribution dominated in smaller size range of aerosol particles with AMAD 0.68 µm (σ_g = 2.18) at height 250 m, and 0.62 µm (σ_g = 2.24) at height 1000 m, showing a dependency on height. In marine environment, the 7Be activity size distribution dominated to higher size range of aerosol particles with AMAD
Table 1. Activity median aerodynamic diameters (AMADs) of \(^7\text{Be}\)-aerosols.

<table>
<thead>
<tr>
<th>(^7\text{Be}) AMADs ((\mu\text{m}))</th>
<th>Latitude</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.76–1.18 (avg. 0.90)</td>
<td>40°38’N</td>
<td>This work</td>
</tr>
<tr>
<td>0.29–0.50 (avg. 0.35)</td>
<td>35°58’N</td>
<td>Bondietti \textit{et al.} (1987)</td>
</tr>
<tr>
<td>0.65–1.09 (avg. 0.77)</td>
<td>51°32’N</td>
<td>Reineking and Porstendorfer (1995)</td>
</tr>
<tr>
<td>0.44–0.74 (avg. 0.57)</td>
<td>48°13’N</td>
<td>Winklet \textit{et al.} (1998)</td>
</tr>
<tr>
<td>1.12–2.06</td>
<td>54°41’N</td>
<td>Lujaniene \textit{et al.} (2001)</td>
</tr>
<tr>
<td>0.33–1.15 (avg. 0.67)</td>
<td>22°18’N</td>
<td>Yu and Lee (2002)</td>
</tr>
<tr>
<td>0.70</td>
<td>51°32’N</td>
<td>Grundel and Porstendorfer (2004)</td>
</tr>
<tr>
<td>0.53</td>
<td>28°04’N</td>
<td>Mohamed (2005)</td>
</tr>
</tbody>
</table>

0.82 \(\mu\text{m} (\sigma_\text{g} = 1.88)\).

Bondietti \textit{et al.} (1987) in thirteen measurements in an almost one-year period at Oak Ridge, Tennessee with temperate latitude (35°58’N, 84°30’W) and wet climate showed that the activity median aerodynamic diameter, AMAD varied from 0.29-0.50 \(\mu\text{m}\) (average 0.35 \(\mu\text{m}\)) and that the fraction of \(^7\text{Be}\)-associated aerosols above 1.4 \(\mu\text{m}\) was usually between 5 and 10%, i.e. analytically, 4.5% was found in the 1.4-2.1 \(\mu\text{m}\) size range, 1.1% in the 2.1-4.2 \(\mu\text{m}\) size range, and only 0.2% in sizes greater than 4.2 \(\mu\text{m}\). They also concluded that cosmogenic radionuclides, such as \(^7\text{Be}\) and \(^{35}\text{S}\) were associated with smaller aerosols in respect of the longer-lived radionuclides, such as \(^{210}\text{Pb}\), a decay product of \(^{222}\text{Rn}\), of terrestrial origin, which were associated with larger-sized aerosols.

Röbig \textit{et al.} (1980) reported that the distribution of \(^7\text{Be}\) is shifted to larger particle sizes due to large residence time of \(^7\text{Be}\) in the atmosphere. An equivalent aerodynamic diameter about 0.65 \(\mu\text{m}\) for \(^7\text{Be}\) might be resulted from the plot of the activity size distribution of the ambient air obtained by a high volume cascade impactor (flow rate 68 m\(^3\)/h) at Göttingen, Germany (51°32’N, 9°55’E). Shifts to large particle sizes were also observed when the relative humidity increased during rainfalls. Very recently, Grundel and Porstendörfer (2004) observed that the results of the \(^7\text{Be}\) measurements for a period of four weeks carried out in outdoor air of a suburb area of the town of Göttingen showed no activity fraction in the nucleation (Aitken nuclei) mode, but a small amount of the activity (5%) in the coarse mode size range (Fig. 4). The accumulation mode of \(^7\text{Be}\)-aerosols with an activity fraction of 95% has an AMAD\(_\text{a}\)-value of 702 nm. The activity size distribution of \(^7\text{Be}\)-aerosols depends probably on the location of formation. Most of \(^7\text{Be}\) atoms and the \(^7\text{Be}\)-aerosols is generated in the upper region of the atmosphere, where other aerosol conditions exist than in the lower atmosphere. Earlier, Reineking and Porstendörfer (1995) by using a Berner-type cascade impactor in an one-year period at Göttingen, Germany, reported AMAD values for the \(^7\text{Be}\)-aerosols in the outdoor atmosphere varying between 650 and 1094 nm (average...
767 nm) and the geometric standard deviation, $\sigma_g$ varied between 1.8 and 2.5 (average 2.1), and that the increase of the AMAD values of $^7$Be-aerosols can in respect of radon decay product aerosols be explained by coagulation processes and the average residence times of about 17-22 days.

Winkler et al. (1998) in forty six measurements in a period of 1 1/3 years at Munich-Neuherberg, Germany (48°13'N, 11°36'E) at a semi-rural area, 490 m above sea level showed that the AMAD of $^7$Be-aerosols ranged from 0.44-0.74 $\mu$m (average 0.57 $\mu$m) and that seasonal effect during the period of high $^7$Be air concentrations, i.e. in the summer, relatively low values of the AMAD (0.45-0.52 $\mu$m) have been observed. They also concluded that the activity median aerodynamic diameter, AMAD ranged between the mass median aerodynamic diameter, MMAD and the surface median aerodynamic diameter, SMAD of the ambient aerosols, indicating that this radionuclide is involved in the transformation process of the tropospheric aerosols after formation in the stratosphere and upper troposphere.

Yu and Lee (2002) in fourteen measurements at Hong Kong (22°18’N, 114°10’E) for a 3½-month period resulted that the activity median aerodynamic diameter of $^7$Be-aerosols varied from 0.33-1.15 $\mu$m (average 0.67 $\mu$m). They concluded that the AMAD of $^7$Be-aerosols is anticorrelated to $^7$Be concentrations in air, is correlated to relative humidity, RH and the mean cloud cover, while temperature does not affect the AMAD of $^7$Be-aerosols.

Mohamed (2005) very recently by using a low-pressure Berner-type cascade impactor at El-Minia, Egypt (28°04’N, 30°45’E) found an average AMAD value of $^7$Be-aerosols in outdoor air 530 nm with relative geometric standard deviation $\sigma_g = 2.4$ and that $^7$Be as a gas diffuses more effectively to a smaller surface area of smaller particles because of the higher surface of these particles.
Apart of this, Lujaniene et al. (2001) reported larger $^7$Be AMAD values varied from 1.12-2.06 $\mu$m (average 1.45 $\mu$m) in a northern latitude (54°41’N, 25°19’E) at Vilnius, Lithuania.

The investigated data (Table 1) rather indicate that the activity median aerodynamic diameter, AMAD of $^7$Be-aerosols increase with increasing latitude (latitudinal effect). As cosmic radiation increases with latitude, the number of $^7$Be atoms and ions formed increase also with latitude, and so there are more $^7$Be-atoms available either to form small aerosol particles in the nucleation (Aitken nuclei) mode and then growing or to be attached directly to the existing large particles in the accumulation mode or in the coarse particle mode by increasing the AMAD of $^7$Be-aerosols.

**Residence Times of $^7$Be Atmospheric Aerosols**

The method for estimating the residence time of atmospheric aerosol particles associated with the radioactive nuclides, such as $^7$Be of cosmogenic origin, is based on the aerosol particle growth rate.

Assuming that the resulting aerosols growth rate (the change in particle diameter, $D_p$ with respect to time $t$) ranged from 0.004-0.005 $\mu$m/h as might be derived from the best fit for droplet phase reactions for the accumulation mode aerosols according to the theory of secondary ambient aerosol growth by condensation and coagulation (McMurry and Wilson, 1982; 1983), then, the residence time, $\tau_R$ of the aerosol can be calculated by dividing the difference between the mean activity median aerodynamic diameter (AMAD)$_{\text{mean}}$ that is of 0.90 $\mu$m for $^7$Be-aerosols (Table 1) and the mean size of Aitken nuclei particles, that is 0.015 $\mu$m (NRC 1979), by the mean particle growth rate, MGR according to the equation

$$\tau_R = \frac{(\text{AMAD})_{\text{mean}} - (\text{AMAD})_{\text{Aitken}}}{\text{MGR}}$$

where MGR is the mean growth rate (0.004-0.005 $\mu$m/h) (McMurry and Wilson, 1982).

Taking into account that the AMAD of aerosol particles associated with $^7$Be varied from 0.76-1.18 $\mu$m (Table 1), then according to the Eq. (2) the residence time of atmospheric aerosols will vary between 7.4 and 8.9 days (average 8.0 days) at Thessaloniki region (40°38’N, 22°58’E), Northern Greece, with dry (precipitation-free) climate at temperate latitude, based in twelve measurements of aerosol samplings carried out during 1½-year period, thus included all seasons of a year.

Papastefanou and Bondietti (1991) reported mean residence times of 8 days for atmospheric aerosols in the boundary layer as determined from $^{210}$Bi/$^{210}$Pb activity ratios at Oak Ridge, Tennessee with temperate latitude (35°58’N, 84°30’W) and wet climate. Therefore, in an attempt to estimate tropospheric aerosol residence times for cosmic-ray spallation products, such as $^7$Be, longer residence times for tropospheric air can be attributed to the influence of stratospheric aerosol contributions as the residence times of
about one week are considered to be valid for tropospheric aerosols at all level of the
troposphere (Martell and Moore, 1974).

Table 2 shows data for the residence times, \( \tau_R \) of atmospheric aerosol particles associate
with \(^{7}\text{Be} \) atoms. The reported from the
literature \( \tau_R \) values for \(^{7}\text{Be}-\text{aerosols} \) varied
from 2.6-35.4 days.

Winkler et al. (1998) estimated residence times 5-6 days in forty six measurements for
\(^{7}\text{Be}-\text{aerosol} \) samplings carried out during 1\( \frac{1}{3} \)-
year period in ground level air at a semi-rural area at Neuherberg, Germany (48°13′,N, 11°36′,E), 490 m above sea level.

Shapiro and Forbes-Resha (1976), much earlier estimated a mean residence tropospheric aerosol residence time for \(^{7}\text{Be-} \)bearing aerosols of 35.4 days, significantly higher, i.e. more than four times higher, at Fullerton, California (33°52′N, 117°55′W), also at mid-latitude for an almost 2-year period with relatively light precipitation.

Yu and Lee (2002) recently estimated mean residence times for \(^{7}\text{Be-associated} \) aerosols ranging from 2.6-11.8 days in fourteen measurements of aerosol samplings carried out during a 3\( \frac{1}{2} \)-month period (November-March), 20 m above ground at Hong Kong, China (22°18′N, 114°10′E) including winter and spring measurements.

Balkanski et al. (1993) following a global three-dimensional model which uses meteorological parameters, such as precipitation scavenging, found that the
tropospheric residence time is a function of latitude (latitudinal effect) according to the
following equation (Ehhalt, 1973).

\[
\tau_R = \frac{C}{\Phi} \tag{3}
\]

where \( C \) is the tropospheric column of a radionuclide extending from the surface up to
the model layer just below the tropopause, and \( \Phi \) is the total depositional flux out of the
column at a given latitude.

Koch et al. (1996) following a three-dimensional chemical tracer model as
Balkanski et al. (1993) also found that the
tropospheric residence time is a function of
latitude according to Eq. (3)

The data of Table 2 admit residence times
of tropospheric aerosols in the range 2.6-35.4
days, but crowd into two groups of values 2.6-
15 days (average 8.8 days) and 21-35.4 days
(average 28.2 days). The lower values are
applicable only to the boundary layer near
Earth’s surface and the higher values are
appropriate to the troposphere as a whole
came to the opposite conclusion, namely, that
the high values are due to the contribution of
stratospheric aerosols, while the lower values
represent the true tropospheric residence time
essentially independent of altitude.

CONCLUSIONS

The aerodynamic size distribution of \(^{7}\text{Be-} \)aerosols in ambient air were measured using normal (1-ACFM) and/or high-volume (HVI)
cascade impactors. Beryllium-7 was largely
associated with submicron aerosol particles in
the accumulation mode (0.4-2.0 µm). Based
on eleven measurements of aerosol samplings,
the activity median aerodynamic diameter,
Table 2. Residence times, $\tau_R$ of tropospheric aerosols.

<table>
<thead>
<tr>
<th>Investigation</th>
<th>$\tau_R$ (days)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thessaloniki, Greece (40°38'N, 22°58'E)</td>
<td>7.4-8.9 (avg. 8.0)</td>
<td>This work</td>
</tr>
<tr>
<td>Neuherberg, Germany (48°13'N, 11°36'E)</td>
<td>5.6</td>
<td>Winkler et al. (1998)</td>
</tr>
<tr>
<td>Fullerton, California (33°52'N, 117°55'W)</td>
<td>35.4</td>
<td>Shapiro and Forbes-Resha (1976)</td>
</tr>
<tr>
<td>Hong Kong, China (22°18'N, 114°10'E)</td>
<td>2.6-11.8</td>
<td>Yu and Lee (2002)</td>
</tr>
</tbody>
</table>

AMAD of $^7$Be-aerosols ranged from 0.76 - 1.18 $\mu$m with a mean value of 0.90 $\mu$m, indicating post-condensation growth either in the upper atmosphere or after mixing into the boundary layer. In the 65% of the 1-ACFM cascade impactor measurements the activity size distribution of $^7$Be-aerosols showed maxima in the 0.7-1.1 $\mu$m size range. The investigated data rather indicate that the activity size distribution of $^7$Be-aerosols increase with increasing latitude (latitudinal effect). Estimated lifetimes of $^7$Be-aerosols in ambient air resulted in a mean residence time of about 8 days that could be applied to aerosol particles in the lower atmosphere below the boundary layer. Contribution of stratospheric aerosols by intrusions could lead to higher values of residence time of atmospheric aerosols, as the residence times of stratospheric aerosols are of a month or higher (Martell, 1970; NRC, 1979; Friedlander, 2000).

NOMENCLATURE

- $1$-ACFM Normal flow rate cascade impactor
- AMAD Activity median aerodynamic diameter
- C Tropospheric column of a radionuclide
- $Cfm$ Cubic feet per meter
- $D_p$ Particle diameter
- $ECD$ Effective cutoff diameter
- $HVIC$ High volume cascade impactor
- $MGR$ Mean particle growth rate
- $MMAD$ Mass median aerodynamic diameter
- $SMAD$ Surface median aerodynamic diameter
- $T_{1/2}$ Radioisotope half-life
- $Z$ Atomic number

GREEK SYMBOLS

- $\lambda$ Radioisotope disintegration rate
- $\sigma_g$ Geometric standard deviation
- $\tau$ Radioisotope mean life
- $\tau_R$ Residence time of aerosol particles
- $\Phi$ Total depositional flux

REFERENCES


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