Beryllium-7 and Lead-210 are Associated with an Increase in the Arctic Oscillation: Evidence from Atmospheric Aerosols in a Remote Tropical Region in East Asia

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Abstract

The Arctic Oscillation (AO) accounts for a large fraction of recent decadal climate trends in Northern Hemisphere (NH) high latitudes. In East Asia, an elevated AO index (AOI) was associated with warmer temperatures in middle- to high-latitude regions, colder temperatures over low-latitude regions, and elevated ozone intrusion from the stratosphere. Elevated beryllium-7 ($^7$Be) is produced in the stratosphere. Few studies have discussed the relationship between $^7$Be and the AO. Here, we identify the AO signature in $^7$Be and lead (Pb)-210 observed at a tropical ambient monitoring site in Nanning (22.8°N, 108.5°E) during the December 2014–December 2017 period. Our results show that the $^7$Be and $^{210}$Pb concentrations are positively and significantly correlated with the AOI ($P < 0.01$). These results show that elevated $^7$Be and $^{210}$Pb are associated with an increase in the AOI, reflecting air masses originating from NH high latitudes and vertically from the high-latitude upper troposphere and lower stratosphere regions to tropical latitudes in East Asia. These results have been verified with ozonesonde data without seasonality and with two meteorological data sets. Our results are also confirmed by observational data over the Pacific regions. We conclude that the AO exerts impacts over the tropical regions in East Asia, and $^7$Be can be used as a tracer to track the impacts of the AO.

Keywords Beryllium-7 · Lead-210 · Arctic oscillation · Atmospheric aerosols · Nanning

1 Introduction

The Arctic Oscillation (AO) is the primary reason for the interannual variability in the troposphere and lower stratosphere. The AO affects the extratropical surface climate variability in the Northern Hemisphere (NH), especially during winter and spring (Thompson and Wallace 2000; Hurrell et al. 2003). Its variability is commonly manifested by a sea-level pressure (SLP) difference between stations in the Azores (Portugal) and Iceland (Thompson and Wallace 1998; Cutlip 2000; Thompson and Wallace 2000; Thompson and Wallace 2001; Thompson and Lorenz 2004). The AO initially emanates from the stratosphere and ultimately alters surface weather, subsequently affecting the temperature and precipitation in most NH areas. Hence, the AO is closely related to the frequency and intensity of extreme weather events (Thompson and Wallace 1998; Thompson and Wallace 2000; Thompson and Wallace 2001; Black 2002; Thompson and Lorenz 2004).

In particular, the AO accounts for a large fraction of recent decadal climate trends in high northern latitudes (Hurrell 1995; Thompson et al. 2000). Thompson et al. (2001) found that in recent decades, the continuous high trend of the North Atlantic Oscillation (NAO, which can be viewed as an AO subset (Jevrejeva and Moore 2001)) is an important reason for winter warming in the NH. In East Asia, as the winter AO index increases from low to high, the temperatures in the middle- and high-latitude regions of the Asian continent...
become warmer, the low-latitude regions become colder, and vice versa (Wu and Wang 2002; Gong et al. 2001; Jeong and Ho 2005; Chen et al. 2013).

Beryllium-7 ($^{7}$Be) is a sensitive indicator of stratospheric air intrusion into the troposphere and this radionuclide is classically applied in the stratosphere-troposphere exchange (STE) (Dibb et al. 1992; Bonasoni et al. 1999, 2000a, b; Cristofanelli et al. 2007, 2006, 2009). In contrast, $^{210}$Pb has been used as a continental tracer of air masses and long-range transport of chemical constituents (derived from continental sources) (Balkanski et al. 1983; Turekian et al. 1983; Baskaran 2011). The cosmogenic radionuclide $^{7}$Be (its half-life is 53.3 days) is produced by high-energy spallation interactions between galactic cosmic-ray (GCR)-produced neutrons and protons (secondary particles) and atmospheric nuclei (primarily nitrogen, oxygen and argon) (Lal and Peters 1967; UNSCEAR 2000; Usoskin and Kovaltsov 2008; Papastefanou 2009; see Fig. S1).

Increased ozone over much of Asia results from the positive phase of the AO combined with changes in the stratosphere-troposphere exchange (STE; Holton et al. 1995; Wang et al. 2002; Wang and Kau 2015) that are responsible for the correlation pattern between ozone and AO at 800 hPa (Lamarque and Hess 2004). This study shows that the tropospheric ozone concentration over Asia is affected by the intrusion of stratospheric ozone. Similar to air containing elevated ozone levels in the stratosphere, an elevated $^{7}$Be level occurs in the stratosphere. Hence, $^{7}$Be is a stratospheric tracer that can be directly or indirectly influenced by the AO.

Similar to $^{7}$Be, ozone from the stratosphere is considered an indicator of the Arctic Oscillation (Thompson and Wallace 2000; Lamarque and Hess 2004; Dibb et al. 1992). However, few studies have discussed the relationship between $^{7}$Be and the Arctic Oscillation. It is worth discussing whether $^{7}$Be can be used as a tracer to track the Arctic Oscillation and whether the signal of the Arctic Oscillation can be discovered through aerosol data in the tropics to study the mechanism of climate teleconnection. Production of $^{7}$Be is the highest in the stratosphere (75%), while the remaining part (25%) is produced in the upper troposphere (Johnson and Viezee 1981; UNSCEAR 2000; Usoskin and Kovaltsov 2008). $^{7}$Be production increases with altitude and geomagnetic latitude and is also associated with the 11-year solar cycle that modulates cosmic-ray penetration through the Earth’s magnetic field (Lal and Peters 1967; Bhandari and Lal 1970; Dibb et al. 1992; UNSCEAR 2000; Usoskin and Kovaltsov 2008). The polar stratosphere experiences the highest production rate, and the tropical lower troposphere exhibits the lowest production rate (Feely et al. 1989).

$^{210}$Pb (with a half-life of 22.3 years) and its progeny polonium, (Po)-210, originate from the naturally occurring radionuclide uranium, (U)-238, in the environment; this can produce radium, (Ra)-226, and radon, (Rn)-222, which are precursors of lead, (Pb)-210 ($^{210}$Pb). Due to the continental origin of $^{222}$Rn, $^{210}$Pb is considered a tracer of air masses with continental origins (Balkanski et al. 1983; Turekian et al. 1983; Baskaran 2011). $^{222}$Rn emanates primarily from rocks and minerals in the crust. Therefore, the spatial variability of $^{210}$Pb is strongly dependent on the geographical types of terrestrial surfaces, and the $^{222}$Rn flux from the ocean is negligible (San Miguel et al. 2019).

$^{210}$Po and $^{210}$Pb nuclides are concentrated in the air over inland cities and industrial areas because of artificial sources, such as industrial mining, automobile exhaust and construction dust (burning of coal, use of phosphate fertilizers, car exhaust, and fires) (Jaworowski et al. 1980; Hotzl and Winkler 1987; Lozano et al. 2013). Once formed in the air, $^{7}$Be, $^{210}$Pb, and $^{210}$Po are rapidly combined with submicron-sized aerosol particles (Maenhaut et al. 1979; Bondietti et al. 1987; Papastefanou 2009). Hence, their concentrations in air are similar to those of aerosol particles and depend on atmospheric transport and wet and dry removal (Papastefanou and Ioannidou 1995; Baskaran 2011).

Because of the different sources of the two radionuclides, simultaneous measurements of $^{7}$Be and $^{210}$Pb and their ratios can be implemented to identify the origin of air masses (Graustein and Turekian 1996; Bonasoni et al. 2000a, b; Zheng et al. 2005) and to study the vertical motion of air masses and convective activity in the troposphere (Brost et al. 1991; Koch et al. 1996; Lee et al. 2004; Tositti et al. 2004; Lee et al. 2007). Moreover, temporal and spatial variations in this ratio reflect both vertical and horizontal transport in the atmosphere (Koch et al. 1996; Baskaran 2011). Due to their half-lives (22.3 years for $^{210}$Pb and 53.2 days for $^{7}$Be), both natural radionuclides can persist long enough for long-range transport in the atmosphere (Zhang et al. 2015; Grossi et al. 2016; San Miguel et al. 2019).

In this paper, we report a three-year (2014/10–2017/12) study conducted on the abundance of $^{7}$Be, $^{210}$Pb, and $^{210}$Po in the surface air over tropical urban site Nanning, China (22.8°N, 108.5°E). In addition, the interrelationship among these nuclides and a suite of related environmental factors, such as the AO, precipitation, cosmic rays, and earthquakes, are also investigated. This paper adapts the Hybrid Single-Particle Lagrangian Integrated Trajectory (HYPLIT) model to calculate the origins of air masses reaching the Nanning sampling site, compares the Nanning results to Environmental Measurements Laboratory (EML) observations, and applies a regression model to study the correlations among the variables of AOI, precipitation, air mass origin (latitude, longitude, and altitude), $^{7}$Be, and $^{210}$Pb.
In this work, we show that air movement in the polar stratosphere affects the tropical troposphere based on observations of radioactive materials near the ground in the tropics. These chemical observations further strengthen the dynamic aspects of stratosphere-troposphere coupling processes (Hoskins et al. 1985). In this paper, we consider the AO as the main mechanism driving air movement variations in the stratosphere-troposphere coupling system. We adopt $^7$Be as a tracer for the stratosphere and $^{210}$Pb as a tracer for the troposphere. With this framework, this work addresses the following issues. (1) A significant correlation has been observed between $^7$Be and AOI, proving that the impact of the AO from

| Code | Start time* | Finish time* | $^{210}$Po (mBq/m$^3$) | $^{210}$Pb (mBq/m$^3$) | $^7$Be (mBq/m$^3$) |
|------|-------------|--------------|------------------------|-----------------------|-------------------|
| 1    | 09.10.2014, / | 10.10.2014, / | 0.55 | 3.92 | 12.7 |
| 2    | 04.11.2014, / | 05.11.2014, / | 0.52 | 1.62 | 9.09 |
| 3    | 08.12.2014,09:19 | 09.12.2014,10:00 | 0.3 | 3.63 | 10.3 |
| 4    | 05.01.2015,10:38 | 06.01.2015,10:39 | 0.48 | 0.71 | 6.08 |
| 5    | 02.02.2015,09:28 | 03.02.2015,09:40 | 0.34 | 3.3 | 1.1 |
| 6    | 04.03.2015,08:25 | 05.03.2015,08:35 | 0.84 | 1.9 | 3.36 |
| 7    | 13.04.2015,10:30 | 14.04.2015,10:30 | 0.9 | 2.13 | 9.37 |
| 8    | 04.05.2015,09:10 | 05.05.2015,09:48 | 0.76 | 0.83 | 1.48 |
| 9    | 16.06.2015,08:58 | 17.06.2015,09:05 | 0.8 | 0.88 | 1.03 |
| 10   | 22.07.2015,09:04 | 23.07.2015,10:18 | 0.59 | 0.36 | 0.9 |
| 11   | 10.08.2015,09:45 | 11.08.2015,09:50 | 0.74 | 0.77 | 1.97 |
| 12   | 23.09.2015,09:27 | 24.09.2015,10:24 | 0.52 | 1.58 | 2.61 |
| 13   | 19.10.2015,10:08 | 20.10.2015,10:23 | 0.69 | 2.43 | 7.56 |
| 14   | 02.11.2015,16:07 | 03.11.2015,16:46 | 0.61 | 2.63 | 4.91 |
| 15   | 16.12.2015,10:51 | 17.12.2015,14:57 | 0.72 | 2.79 | 7.83 |
| 16   | 19.01.2016,10:39 | 20.01.2016,11:19 | 1.19 | 3.12 | 8.41 |
| 17   | 16.02.2016,10:32 | 17.02.2016,10:45 | 0.43 | 1.26 | 8.5 |
| 18   | 01.03.2016,10:05 | 02.03.2016,16:07 | 1.16 | 1.67 | 8.98 |
| 19   | 06.04.2016,14:50 | 07.04.2016,15:07 | 0.34 | 0.92 | 4.66 |
| 20   | 03.05.2016,10:13 | 04.05.2016,10:35 | 0.24 | 2.47 | 3.47 |
| 21   | 27.06.2016,09:50 | 28.06.2016,10:11 | 0.19 | 0.23 | 0.84 |
| 22   | 18.07.2016,08:40 | 19.07.2016,08:52 | 0.31 | 0.34 | 2.47 |
| 23   | 08.08.2016,11:35 | 09.08.2016,11:12 | 0.8 | 0.83 | 1.1 |
| 24   | 18.09.2016,16:00 | 19.09.2016,16:20 | 0.67 | 1.28 | 3.55 |
| 25   | 09.10.2016,09:01 | 10.10.2016,08:56 | 1.87 | 2.65 | 3.52 |
| 26   | 14.11.2016,11:10 | 15.11.2016,11:35 | 0.18 | 0.61 | 0.78 |
| 27   | 07.12.2016,10:42 | 08.12.2016,10:32 | 1 | 1.62 | 9.66 |
| 28   | 03.01.2017,11:50 | 04.01.2017,12:05 | 0.869 | 2.309 | 5.76 |
| 29   | 13.02.2017,10:38 | 13.02.2017,10:38 | 0.515 | 1.412 | 6.55 |
| 30   | 02.03.2017,15:52 | 03.03.2017,15:08 | 1.679 | 2.622 | 7.69 |
| 31   | 17.04.2017,10:56 | 18.04.2017,11:24 | 0.393 | 0.854 | 1.21 |
| 32   | 03.05.2017,17:15 | 04.05.2017,17:22 | 0.223 | 1.129 | 2.14 |
| 33   | 20.06.2017,11:00 | 21.06.2017,09:45 | 0.625 | 0.669 | 0.484 |
| 34   | 20.07.2017,09:45 | 21.07.2017,09:15 | 0.227 | 0.238 | 0.765 |
| 35   | 07.08.2017,11:25 | 08.08.2017,11:16 | 0.517 | 0.372 | 0.743 |
| 36   | 29.09.2017,16:35 | 30.09.2017,17:15 | 0.235 | 0.265 | 0.387 |
| 37   | 30.10.2017,09:00 | 31.10.2017,17:05 | 0.905 | 1.314 | 0.781 |
| 38   | 17.11.2017,16:10 | 19.11.2017,11:50 | 0.172 | 0.298 | 0.649 |
| 39   | 09.12.2017,20:12 | 10.12.2017,19:33 | 0.538 | 2.054 | 0.646 |

*The local time of Nanning is given in the following format: DDMMYY, HH:MM (day, month, year, hour, minute; 08.12.2014 is read as 8 December 2014)
the NH polar stratosphere can be observed near the surface of the tropical troposphere. (2) The good linear correlations between $^7$Be and $^{210}$Pb indicate that the stratosphere and troposphere are coupled.

2 Data and Methods

2.1 Observational Data

2.1.1 Site Description

The sampling site is an automatic environmental station in Nanning city, China (22.8°N, 108.5°E). Nanning has a humid subtropical monsoon climate with an average temperature of 21.7 °C and abundant rainfall (Deng et al. 2018). The average annual rainfall is 1298 mm, and Nanning has a humid summer and a slightly dry winter with distinctive dry and wet seasons (which result from the impact of the Asian monsoon).

2.1.2 Sample Collection

Aerosol samples are collected with a PM-800 large-flow aerosol sampler with a mounted HB1 polypropylene fiber filter membrane and an electric tablet press (Fig. 1). The collection efficiency of the HB1 filter membrane for 0.5 μm aerosol particles is approximately 99%. The operational flow of the sampler is set to 600 m$^3$/h to ensure that the standard-condition volume of sampling is not smaller than 10,000 m$^3$. After sampling, the volume of air and the start and end times are recorded. The sampling frequency is one sample over approximately 24 h every month from October 2014 to December 2017. The total number of samples is 39. Each aerosol sample is compressed into a cylinder with a diameter of 70 mm and a length of 10 mm (Bourcier et al. 2011; Zhang et al. 2015).

2.1.3 Sample Analysis

After encapsulation, the radioactivity was measured with a high-purity germanium gamma (HPGe-$\gamma$) spectrometer system containing a model BE3830 detector (CANBERRA) with a crystal size of $\Phi$ 80 mm × 30 mm. The counting error for radioactivity measurements was approximately 10% at the 1 sigma level (Lee et al. 2004). The linear energy scale of the analyzer was calibrated with photon peaks from a Co-60 source (1332 keV) with the full width at half maximum (FWHM) of energy no higher than 1.80 keV and with a 50.9% relative efficiency. The measurement time was longer than 86,400 s, and the minimum detection limits for $^7$Be, $^{210}$Pb, and $^{210}$Po were 0.017–0.022 mBq/m$^3$, 0.01–0.03 mBq/m$^3$ and 0.001–0.005 mBq/m$^3$, respectively. The uncertainties of $^{210}$Pb and $^7$Be activities were controlled to below 5%, and the uncertainties of $^{210}$Po were approximately 13%.

In this paper, the $^7$Be and $^{210}$Pb activities were determined with $\gamma$ rays of 477.6 keV ($P_\gamma$ = 10.43%) and 46.5 keV ($P_\gamma$ = 4.05%), respectively. The detection efficiency was obtained with laboratory source calibration software (LabSOCS). LabSOCS was considered reliable for gamma ray detector efficiency calculations, and the deviation of detection efficiency calculated by this method was less than 10% (Bronson 2003; Li and Geng, 2010; Done et al. 2016). The $\gamma$ energy spectrum data were analyzed with Genie-2000 spectrum analysis software to obtain the activities and uncertainties of $^{210}$Pb and $^7$Be in the aerosol samples (mBq/m$^3$) (Rastogi and Sarin 2008; Baskaran and Shaw 2001; Leppänen et al. 2010; Dueñas et al. 2011; Du et al. 2020).

After gamma spectrometry analysis, each sample was spiked with $^{209}$Po as a tracer, and the polonium in the sample was then dissolved with concentrated nitric acid. The residue was dissolved/leached with hydrochloric acid, and ascorbic acid was added thereafter. In the hydrochloric acid system, polonium was self-plated onto a silver disc and determined with an alpha spectrometry (Canberra, 7200–4 PIPS) instrument connected to an Alpha Analyst to calculate the activity of $^{210}$Po (Carvalho 1995; Huh and Su 1999; Baskaran and Shaw 2001; Daish et al. 2005; Matthews et al. 2007).

2.2 The HYSPLIT Model

In this work, we apply the National Oceanic and Atmospheric Administration (NOAA) HYSPLIT model (Stein et al. 2016)
to produce back trajectories in the filtering sampling periods with a duration of 168 h (7 days). Back trajectories are calculated every six hours over a specific sampling period. The source site is located in Nanning (latitude: 22.8°N; longitude: 108.5°E), and the height is 100 m above ground level because the air below 100 m can be regarded as a uniform mixing layer. The vertical motion calculation method relies on a vertical velocity model, for which we use the meteorology over the sampling period from the National Centers for Environmental Prediction (NCEP) GDAS0p5 (Global Data Assimilation System 0.5 degrees longitude-latitude). Since the number of samples is 39 and the duration of each sampling period is longer than 20 h, there are a total of 144 back trajectories.

2.3 Linear Regression Search

For a surface-observed variable \(X\), we can define \(X = X(\varphi, \lambda, h, H, P, A, C)\). Here, \(\varphi, \lambda, h\) and \(H\) are determined from the HYSPLIT back trajectories, \(\varphi\) indicates the maximum and minimum latitudes, \(\lambda\) gives the maximum and minimum longitudes, \(h\) represents the maximum and minimum altitudes, \(H\) is the mixing depth of the planetary boundary layer, and \(P\) is the 24-h precipitation rate obtained from the United Kingdom Meteorological Office (UKMO) Hadley Centre. The data analyzed in this work are HadISD version 3.1.0.201911p (Dunn et al. 2012, 2014, 2016). \(A\) is the AOI, which is the daily mean data obtained from the National Centers for Environmental Prediction (NCEP)/National Center for Atmospheric Research (NCAR) reanalysis (Zhou et al. 2001), and \(C\) indicates the ground-level cosmic ray observations obtained from the cosmic ray station of the Sodankyla Geophysical Observatory, University of Oulu, Finland (Mishev et al. 2014).

The daily AO index is obtained from the National Oceanic and Atmospheric Administration (NOAA) Climate Prediction Center (ftp://ftp.cpc.ncep.noaa.gov/cwlinks/norm.daily.ao.index.b500101.current.ascii). The daily AO index is calculated via methods similar to Thompson and Wallace (2000) and Thompson et al. (2000). The daily AO index is determined by projecting the daily 1000 hPa height anomalies at latitudes north of 20°N onto the leading empirical orthogonal function (EOF) (https://www.cpc.ncep.noaa.gov/products/precip/CWlink/daily_ao_index/ao.shtml). The monthly mean 1000 hPa height anomalies north of 20°N are used to construct the EOF. The daily AO index is normalized by the standard deviation of the monthly index (based on the 1979–2000 period). NCEP/NCAR reanalysis data at 2.5 degree longitude-latitude resolution are used in the construction of the monthly EOF (https://www.cpc.ncep.noaa.gov/products/precip/CWLink/daily_ao_index/history/method.shtml). The positive AO index corresponds to negative height anomalies over the polar region, and vice versa (Thompson et al. 2000).

To rank the effects of the above parameters on the surface-observed \(X\), we apply a linear regression model to compute the statistical correlations between both the observed \(X\) and variables and among the variables (Wang and Chau 2013). The correlations between two variables are measured by the calculated correlation coefficient \(R\). The statistical significance of a correlation is tested by computing Student’s t test probability \(P\) (Press et al. 1992).

2.4 The US EML Global Air Sampling Data

The US EML has a network of air sampling sites that measure \(^{7}\text{Be}\), \(^{210}\text{Pb}\), and other nuclides on a global scale (Larsen et al. 1995). The global coverage of EML surface sites is excellent for comparing and testing the results obtained from the analysis of local measurements performed in Nanning. It is important to test whether the results obtained from Nanning are specific only to Nanning or whether they are general features on a global scale. In this context, the EML data serve as a testbed for verifying the results obtained from the Nanning data.

3 Results

3.1 Time-Series Measurements

Time-series measurements of \(^{7}\text{Be}\), \(^{210}\text{Pb}\), and \(^{210}\text{Po}\) over 39 months from 2014 to 2017 are shown in Fig. 2(a). The average specific activity of \(^{7}\text{Be}\) is 4.21 mBq/m³ and is in the range of 0.39–12.70 mBq/m³. The average specific activity of \(^{210}\text{Pb}\) is 1.51 mBq/m³ and is in the range of 0.23–3.92 mBq/m³. The average specific activity of \(^{210}\text{Po}\) is 0.61 mBq/m³ and is in the range of 0.17–1.68 mBq/m³. The activity level of \(^{7}\text{Be}\) is higher than the global average level of 2.45 mBq/m³ based on long-term observations of more than 70 stations around the world (Koch et al. 1996). The activity of \(^{7}\text{Be}\) is comparable to that determined at other observation stations (EML data, as shown in Fig. 5). The activities of \(^{210}\text{Pb}\) and \(^{210}\text{Po}\) in the ambient aerosols mostly exceed the simulated values at 0.5 and 0.05 mBq/m³, respectively (UNSCEAR 2000). However, the activity levels of \(^{210}\text{Pb}\) and \(^{210}\text{Po}\) at Nanning are comparable to those over other cities in China (Pan et al. 2017). The three nuclides exhibit a seasonal variation pattern: higher activity in winter and lower activity in summer. The overall concentration of \(^{7}\text{Be}\) is higher than that of \(^{210}\text{Pb}\) and \(^{210}\text{Po}\), and its fluctuation is also more notable. Rainfall is the main process of aerosol removal from the atmosphere. Rainfall is abundant in the summertime in Nanning, which decreases \(^{210}\text{Pb}\) and \(^{210}\text{Po}\), similar to \(^{7}\text{Be}\). Although the production rate of \(^{7}\text{Be}\) at any given latitude in the atmosphere does not change with the season, the seasonal intrusion of lower stratospheric air and washout of atmospheric aerosols carrying \(^{7}\text{Be}\) leads to the enrichment of \(^{7}\text{Be}\) during late winter and early spring,
which does not occur in summer (Marenco and Fontan 1974; Feely et al. 1989; Dueñas et al. 2020).

$^7$Be is a cosmogenic radionuclide that is mainly produced in the stratosphere. Due to cosmic rays, $^7$Be is continuously produced. Ground-level cosmic ray observations revealed a persistent increasing trend from October 2014 to December 2017. Despite increasing cosmic rays, the activity of $^7$Be in the near-surface air did not persistently increase. Hence, the variations in ground-level $^7$Be concentrations are more associated with AO patterns than with changes in the incoming cosmic rays. The back trajectory analysis (Fig. 2(b)) clearly shows that almost all of the higher concentrations of $^7$Be originate from middle and high latitudes. Some of the trajectories start in polar regions. These back trajectories indicate that $^7$Be observed near the ground in Nanning was affected by air from middle and high northern latitudes.

The activities of $^{210}$Pb and its progeny $^{210}$Po exhibit seasonal variation, with low concentrations in summer and high concentrations in winter. The back trajectory maps of $^{210}$Pb and $^{210}$Po are shown in Fig. 2(b). The back trajectories of the higher concentrations are overwhelmingly distributed over the land area. Oceanic air masses from the South China Sea play a dominant role in affecting the weather in Nanning in summer with negligible $^{222}$Rn activity, while the back trajectories of the land air masses in winter indicate higher $^{222}$Rn activity. Therefore, the activity of the $^{222}$Rn, $^{210}$Pb and $^{210}$Po progenies exhibits the same seasonal distribution in aerosols (Carvalho 1995; Dueñas et al. 2011).

### 3.2 Correlation Analysis

Figure 3(a) shows that the correlation between $^7$Be and the AOI is the most significant, with an R = 0.26, and the P value is 0.0268, followed by $^{210}$Pb, with an R = 0.20 and a P value of 0.0826. The correlation between $^{210}$Po and the AO is an R value of 0.29 and a P value of 0.0138. More noteworthy discussions are presented below. As a tracer of the STE, the correlation of $^7$Be versus $^{210}$Pb is significant, with R = 0.62 and a P value < 0.01 (Fig. 3(c)). Precipitation is negatively correlated with the $^7$Be concentration, with R = 0.30 and a P value = 0.11 (Fig. 3(b)). Note that Fig. 3(a) also shows that low levels of $^7$Be (less than 1 mBq/m$^2$) are also associated with a high AOI, indicating the involvement of multiple processes in the scatter diagram.

Figure 3(b) shows negative correlations between $^7$Be, $^{210}$Pb and precipitation. The negative correlation between $^{210}$Pb and precipitation is characterized by R = 0.35 and a P value of 0.06, while $^{210}$Po shows no clear correlation with precipitation. An earthquake is the source event for $^{210}$Pb and $^{210}$Po. Due to crustal rupture, a high concentration of $^{222}$Rn can diffuse from crustal soil to the atmosphere. Therefore, seismic events may be able to explain the short-term and sudden high-activity events of $^{210}$Pb and $^{210}$Po as long half-life progenies of $^{222}$Rn. However, seismic signals are relatively complex and are not explained in this work.

To clarify the effect of the AO on $^7$Be, we perform a series of sensitivity tests with various durations (from 2 to 7 days) of the back trajectories. The test results reveal that the 6–7 day back trajectories result in the most significant correlation (R = 0.46 and P value < 0.01) between $^7$Be and height. Hence, correlation analysis is performed on these four variables (AOI, $^7$Be, latitude, and altitude) by considering three AOI cases (all, positive phase, and negative phase (see Fig. 4)). For data reliability, a comparison between the EML and Nanning observation data is conducted via scatter plot analysis of $^7$Be versus AOI and of $^{210}$Pb versus $^7$Be.

Figure 4(a) shows that 63% (90/142) of the back trajectories in the case of AOI > 0 originate in the NH polar region and mid-latitude regions. The back trajectories for AOI < 0 (37%)
are confined to latitudes south of 50°N. Both 7Be and the AOI exhibit a significant correlation with the maximum altitudes and latitudes of the HYSPLIT back trajectories (Fig. 4(b, c, d, e)). Especially in the case of AO > 0, all the correlations are more notable. Regarding the AOI versus latitude, the R = 0.28 in the case of all AOI data (P value <0.01; Fig. 4(b), left panel). The correlation coefficient R increased to R = 0.48 in the case of AOI > 0 (P value <0.01; Fig. 4(b), middle panel). In terms of the AOI versus altitude, the R = 0.10 in the case of all AOI data (Fig. 4(c), left panel). The correlation coefficient R increased to R = 0.32 in the case of AO > 0 (P value <0.01; Fig. 4(c), middle panel).

Fig. 3 Scatter plots of the observed variables versus the AOI. a The three radionuclides (7Be (left panel), 210Pb (central panel), and 210Po (right panel)) versus the AOI. b The three radionuclides (7Be (left panel), 210Pb (central panel), and 210Po (right panel)) versus precipitation. c Correlations between 210Pb and 7Be (left panel) and between 210Po and 7Be (right panel)
The correlation between altitude and $^7$Be is significant, with $R = 0.40$ in the case of all AOI data (Fig. 4(d), left panel). The correlation coefficient is $R = 0.37$ in the case of AOI $> 0$ (P value $< 0.01$; Fig. 4(d), middle panel). These results prove that the sources of the elevated $^7$Be are located at high altitudes. The correlations between latitude and $^7$Be ($R = 0.56$ in

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Fig. 4 Scatter plots of $^7$Be measurements versus the highest altitudes and latitudes of the air mass back trajectories from HYSPLIT. a Spatial distribution of the back trajectories: all AOIs (left panel), positive AOIs (middle panel), and negative AOIs (right panel). b All AOIs (left panel), positive AOIs (central panel), and negative AOIs (right panel) versus altitude. c All AOIs (left panel), positive AOIs (central panel), and negative AOIs (right panel) versus latitude. e All AOIs (left panel), positive AOIs (central panel), and negative AOIs (right panel) for positive AOIs.
the case of AOI > 0, with a P value < 0.01; Fig. 4(e), left panel) prove that elevated $^{7}$Be is associated with higher latitudes in the NH. The latitude versus altitude correlations (R = 0.61, with a P value < 0.01; Fig. 4(e), right panel) demonstrate that the high-latitude air masses originated from the NH high latitudes.

The polar stratosphere exhibits the highest $^{7}$Be production rate, while the tropical lower troposphere experiences the lowest $^{7}$Be production rate (Feely et al. 1989). The variations in the AO exhibit chains of coupled flow pattern changes from the polar stratosphere downward toward the lower-latitude troposphere. With the AO as a key mechanism, air parcels move from higher latitudes and altitudes (the polar stratosphere that is rich in $^{7}$Be) to lower latitudes and altitudes (the tropical troposphere) when the AO is positive, and East Asia experiences a more notable STE. Therefore, it is easy to understand that with the higher altitudes and latitudes of the HYSPLIT trajectories, it is more likely for the air enriched in $^{7}$Be to be transported from the stratosphere to troposphere, as well as from the high to low latitudes. These processes explain why the AOI exhibits good correlations with $^{7}$Be. These four aspects ($^{7}$Be, AOI, latitude, and altitude) are notably correlated through analysis of the back trajectories.

In summary, Fig. 4 demonstrates that elevated $^{7}$Be is associated with air coming from high latitudes (polar regions) and high altitudes (lower stratosphere). In the case of AOI > 0, strong and positive regression correlations exist between latitude and altitude versus $^{7}$Be and between altitude versus latitude. In the case of AOI < 0, a weak positive regression is observed. All correlation coefficients are statistically significant (P value < 0.01 for positive phases of the AO). These results indicate that the air masses enriched in $^{7}$Be originate from polar regions and the lower stratosphere. The disturbance and variability of the polar vortex, exhibited as the AOI, are the main causes of the elevated $^{7}$Be observed at the tropical Nanning site.

The back-trajectory computation strongly supports our findings of the dominant role of the AO in controlling the
abundance of ⁷Be (Fig. 3(a)) and the significant correlation between ⁷Be and ²¹⁰Pb (Fig. 3(c)). Moreover, based on the meteorology along the back trajectories, we perform further correlation analyses (Fig. 4) and prove that the air masses enriched in ⁷Be originate from the polar regions and the lower stratosphere.

3.3 Comparisons with EML Data

For data reliability and to test our results, scatter plot analysis of ⁷Be versus AOI and ²¹⁰Pb versus ⁷Be was again performed on the EML data, which contains more than 6000 observations on a global scale. Figure 5 shows the sites in the NH that have similar patterns as Nanning. Figures 6 and 7 show the sites in the Southern Hemisphere (SH) that exhibit both positive correlations between the AOI and ⁷Be and between ⁷Be and ²¹⁰Pb. The correlation analysis results are statistically significant.

In Perth (32°S), Australia, ⁷Be is correlated with the AOI with an R = 0.11, and ²¹⁰Pb is correlated with ⁷Be with an R = 0.36 (P < 0.01). In Tasmania (40.73°S), Australia, the ⁷Be versus AOI correlation has an R-value of 0.11, and the ²¹⁰Pb versus ⁷Be correlation has an R-value of 0.21 (P < 0.01). On Reunion Island (21.10°S), the ²¹⁰Pb versus ⁷Be correlation has an R-value of 0.36 (P < 0.01). The comparisons show that similar latitudes exhibit similar ranges of ⁷Be concentrations and correlation coefficients between ²¹⁰Pb and ⁷Be (R = 0.31, P < 0.01), while the ²¹⁰Pb correlation coefficient range is half of that at the Nanning site (due to abundant fossil fuel burning). The patterns from the 5 Australian sites are consistent, which verifies that they are correct.
advection and the disturbance between the troposphere and stratosphere. As a tracer of the STE, the good coherence between $^7$Be and $^{210}$Pb indicates that the disturbance of the air due to high mountains can promote the coupling of the
stratosphere and troposphere. Interestingly, Nanning exhibited correlations of both $^7$Be versus AOI and $^{210}$Pb versus $^7$Be that were higher than those at all the EML sites. Nanning is located in the westerly disturbance area on the leeward side of
the Himalayas, which is the highest mountain range on Earth. Hawaii, affected by the Himalayas, is downwind of Nanning. Montgomery, Alabama, is on the east coast of the US and experiences a relatively severe air disturbance due to the Rocky Mountains. In Barrow, Alaska, which is affected by the Alaska Mountain Range, the correlations between $^{210}$Pb and $^7$Be are also notable with $R = 0.59$ and a $P$ value $<0.01$, and the lower $^{210}$Pb activity may be due to its coastal location. The Montgomery site (32°N) in Alabama exhibits a good correlation between $^{210}$Pb and $^7$Be, with $R = 0.49$ ($P < 0.01$).

Figure 8(b) shows the sites that exhibit statistical significance in the linear regression analysis of $^7$Be versus $^{210}$Pb. This figure shows the sites that are the most responsive and reflect the highest degree of coupling between the stratosphere and troposphere during the positive phases of the AO.

At these sites, the correlations between $^7$Be and AOI and between $^{210}$Pb and $^7$Be are significant with $P$-values that are less than 0.05; these sites include Barrow, Alaska and Montgomery, Alabama in the USA, and Perth and Tasmania in Australia. Note that Fig. 7(e) shows that both Nanning and Mawson Station in Antarctica have similar measurements of $^7$Be, but the $^{210}$Pb at Mawson is very low compared with that at Nanning. The higher $^{210}$Pb activity at the Nanning site is mainly due to its location in an inland city as well as industrial influence. Mawson Station in Antarctica, which is far from continents, has very low $^{210}$Pb activity due to the reduction (decay and diffusion) in $^{210}$Pb during long-distance ground dust transport from remote terrestrial areas. On the other hand, the elevated $^7$Be over Mawson, Antarctica indicates the abundant sources of $^7$Be from the lower stratosphere over Antarctica.

The good correlations between $^7$Be and $^{210}$Pb represent the spatial distribution of observations dominated by stratosphere-troposphere coupled effects (Fig. 8(b)). The good correlations between the AOI and $^7$Be represent the spatial distribution of sites with the AO-initiated stratospheric effect in the troposphere (Fig. 8(a)). The combined good correlations of the paired AOI versus $^7$Be and $^7$Be versus $^{210}$Pb represent the AO-initialized stratosphere-troposphere coupled effect.

4 Discussion and Conclusions

In this work, we showed that the ground levels of $^7$Be and $^{210}$Pb at the Nanning and EML stations in Alaska (Barrow), Montgomery (Alabama), Australia (Perth, Tasmania, and Norfolk), New Zealand (Chatham), and American Samoa were positively and statistically significantly correlated with an increase in the positive AOI. What are the mechanisms resulting in the increase in $^7$Be and $^{210}$Pb being positively correlated with the AOI? What processes that lead to $^7$Be were positively correlated with $^{210}$Pb at these sites?

The production rates of $^7$Be are higher in the upper troposphere and lower stratosphere (25–50 $^7$Be atoms s$^{-1}$ (gram of air)$^{-1}$) than in the lower troposphere (1–5 $^7$Be atoms s$^{-1}$ (gram of air)$^{-1}$; Koch et al. 1996). Approximately two-thirds of the $^7$Be is produced in the stratosphere, and one-third of the $^7$Be is produced in the troposphere (Rehfeld and Heimann 1995). On the other hand, $^{210}$Pb is produced from the radioactive decay of $^{222}$Rn, which is released from terrestrial soil (Koch et al. 1996). Hence, only a small contribution of stratospheric air is needed to produce a large change in the observed $^7$Be in surface air (Dutkiewicz and Husain, 1985).

Figure 9 shows the difference in geopotential height (m) at 850 hPa between years of positive AO (1989, 1990, 1992, 1993, 1997, and 2000) and years of negative AO (1980, 1985, 1987, and 2001) calculated by Lamarque and Hess (2004). The positive geopotential anomalies are associated with the positive anomalies of high pressures and clockwise circulations. Red curves encircle the regions of the anomalous highs with high pressure centers highlighted by $H_1$, $H_2$, $H_3$, and $H_4$. $H_1$ is centered at approximately 120°E and 45°N (eastern Asian continent). $H_2$ is centered around the North Pacific, at 150°W and 60°N. The high-pressure systems are
accompanied by subsidence flow from the upper troposphere to the lower stratosphere. The lower panel of Fig. 9 shows vertical circulations associated with the anomalous high- and low-pressure systems. Hence, high-pressure systems are associated with enhanced subsidence (Lee et al. 2004). They are conducive to the vertical transport of elevated $^7$Be from the upper troposphere and lower stratosphere (UTLS) regions to the lower troposphere. The subsidence associated with high-pressure systems also suppresses convection and precipitation processes in the atmospheric boundary layer, leading to the enhancement of $^{210}$Pb at the surface (Rehfeld and Heimann 1995; Koch et al. 1996; Wang et al. 1999). Hence, the development of the positive AOI is conducive to the transport of elevated $^7$Be from the UTLS regions to the lower troposphere.

The negative differences in the AOI are associated with the negative anomalies of the low pressure and anti-clockwise circulations. They are associated with enhanced upward motion. The development of the anomalous low-pressure system, L, over the Arctic region acts to enhance the strength of the polar vortex (Thompson and Wallace 1998; Baldwin and Dunkerton 1999; Ambaum and Hoskins 2002). A developing polar vortex produces atmospheric circulations similar to those of the AO patterns in Fig. 9 and is conducive for the vertical transport of elevated $^7$Be and downward transport of elevated potential vorticity (PV) and $^7$Be to the lower troposphere (Black 2002; James et al. 2003).

The Nanning site is located downstream of the $H_1$ high-pressure system, while the $H_2$ high-pressure system influences the Barrow site in Alaska during the positive AO phases. The rest of the sites (Mauna Loa, three sites in Australia, two sites in New Zealand, and American Samoa) are located in regions with enhanced subsidence during the positive AO phases (Limpasuvan et al. 2005; Zhao and Miller 2005).

The HYSPLIT model calculations show that the back trajectories tend to originate in the high altitudes of the troposphere and with the origins of the trajectories in northern higher latitudes. These results are presented in the regression model analysis (altitudes are correlated with latitudes). The clockwise back trajectories are consistent with the subsidence of the flows associated with the high-pressure systems from northern high latitudes and high altitudes to the tropical Nanning site.

We used NCEP reanalysis data to check the sensitivity of the HYSPLIT results with respect to the meteorological data used in this work. Fig. S2 compares the horizontal distribution of the back trajectories calculated with the GDAS0p5 data (Fig. S2(a)) and with the NCEP reanalysis data (Fig. S2(b)). We note that the GDAS0p5 data have a horizontal resolution of 0.5 degrees longitude-latitude and contain output data collected every 3 h on 55 hybrid sigma-pressure levels from the surface to 13 hPa. The reanalysis data have a horizontal resolution of a 2.5 degree longitude-latitude grid, output data collected every 6 h, and 17 pressure levels (1000, 925, 850, 700, 600, 500, 400, 300, 250, 200, 150, 100, 70, 50, 30, 20, and 10 hPa). Fig. S2 shows that the general patterns of the gridded distribution of the occurrence of back trajectories are similar. However, the results calculated with GDAS0p5 data show a more widespread distribution of the back trajectories than the results calculated with the reanalysis data. The GDAS0p5 data have a higher horizontal and vertical resolution than the reanalysis data, resulting in a better representation of the vertical velocity. The linear regression analysis of the AOI and $^7$Be versus latitudes and altitudes calculated with the reanalysis data are consistent with the results calculated with the GDAS0p5 data. Elevated AOs are associated with air originating from northern higher latitudes (Fig. S3(a)). Elevated $^7$Be is associated with air originating from northern higher latitudes (Fig. S3(b)). The air from northern higher latitudes originates from elevated altitudes (Fig. S3(c)). Elevated AOs are associated with air from higher altitudes (Fig. S3(d)). Elevated $^7$Be concentrations are associated with air originating from higher altitudes (Fig. S3(e)). The altitudinal origins of the air calculated with the reanalysis data are mostly limited to altitudes below 5 km (Fig. S3(d)), which are lower than the origins of the air calculated with the GDAS0p5 data (Fig. 4(d)). These comparisons indicate that the vertical velocities

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**Fig. 9** A schematic diagram showing the distribution of the anomalous high- and low-pressure systems associated with differences between the positive and negative AO years. The shaded colors indicate the distribution of the anomalous high (red) and low (blue) pressure systems adapted from Lamarque and Hess (2004). The locations of the high-pressure systems are indicated by $H_1$, $H_2$, $H_3$, and $H_4$. Schematic flow symbols indicate that the main low-pressure system, L, over the Arctic regions is associated with the anomalous high- and low-pressure systems. The bottom panel shows a meridional vertical circulation through the 120°E and 60°W cross-sections (indicated by the green line in the upper panel) associated with the anomalous high- and low-pressure systems.
Beryllium-7 and Lead-210 are Associated with an Increase in the Arctic Oscillation: Evidence from...

...and Lead-210 are associated with an increase in the Arctic Oscillation: Evidence from...(19.43°N, 155.04°W; Fig. S 4(g)). In contrast, decreases in H2O are associated with increases in the AOI at Taipei (22.31°N, 114.17°E; Fig. S 4(d)), and Hilo, Hawaii. O3, H2O, and temperature versus AOI at three sites over the 2021). Fig. S 4 shows the linear regression model analysis of and temperature from ozonesonde measurements (WOUDC shown in this study, the levels of near-surface 7Be and 210Pb are since 1950 (e.g., Visbeck et al. 2001; Hurrel et al. 2004). As the Arctic Oscillation (NAO) and AO have become more positively biased over North Pacific flight corridors can be traced back to the ground level industrial areas over East Asia. In contrast, the back trajectories show that low CO in the upper troposphere is associated only with the air in the upper troposphere, where the CO concentrations are low.

We note that Nanning has abundant precipitation leading to wet summers, and rainfall is the main removal process of aerosols from the atmosphere. Therefore, the influence of removal of rainfall on radionuclides in aerosols should be taken into account. Therefore, in Fig. 3(b), we considered the correlation analysis of rainfall on the activity of the three nuclides. The negative correlations shown in Fig. 3(b) are consistent with previous works (e.g., Rehfeld and Heimann 1995; Koch et al. 1996), showing the wet scavenging effect of 7Be and 210Pb in Nanning.

The dynamics of recent climate change have been linked to the Arctic (e.g., James et al. 2003). The North Atlantic Oscillation (NAO) and AO have become more positively biased since 1950 (e.g., Visbeck et al. 2001; Hurrel et al. 2004). As shown in this study, the levels of near-surface 7Be and 210Pb are positively correlated with the increasing trends in the positive AO. Continuous measurements of near-surface 7Be and 210Pb can provide radioactive evidence for monitoring continuous changes in the AO and the links to global warming.

Based on previous works (Wang et al. 2002; Wang and Kau 2015 and references therein; and references cited in the manuscript), we use the association between ground-level observed 7Be (a stratospheric tracer) and the AOI to indicate the changes in the STE associated with the AOI in this work. The air with stratospheric origins contains elevated 7Be, elevated O3, low CO, and low H2O. Hence, we also analyze O3, H2O, and temperature from ozonesonde measurements (WOUDC 2021). Fig. S 4 shows the linear regression model analysis of O3, H2O, and temperature versus AOI at three sites over the North Pacific from 2014 to 2017. At an altitude of 3 km (to reduce the impact of ground-level photochemical O3 production), increases in O3 are associated with increases in the AOI at Taipei (25.00°N, 121.44°E; Fig. S 4(a)), Hong Kong (22.31°N, 114.17°E; Fig. S 4(d)), and Hilo, Hawaii (19.43°N, 155.04°W; Fig. S 4(g)). In contrast, decreases in H2O are associated with increases in the AOI at Taipei (Fig. S 4(b)) and Hong Kong (Fig. S 4(e)). H2O at Hilo (Fig. S 4(h)) was compounded by H2O evaporation from the underlying ocean. Additionally, decreases in AOIs are associated with decreases in temperature in Taipei (Fig. S 4(c)), Hong Kong (Fig. S 4(f)), and Hilo (Fig. S 4(i)). Fig. S 5 shows regression analysis profiles throughout the troposphere, from the surface to an altitude of 10 km, with respect to increases in the AOI. At Taipei, O3 increases at altitudes of 3 km and above 5 km (Fig. S 5(a)), and decreases in H2O (Fig. S 5(b)) and temperature (Fig. S 5(c)) occur in the troposphere. At Hong Kong, O3 increases at altitudes below 7 km (Fig. S 5(d)), and decreases in H2O (Fig. S 5(e)) and temperature (Fig. S 5(f)) occur in the troposphere. Hong Kong is located 580 km east of the Nanning site. Hong Kong has ozonesonde data that is closest to the Nanning site. In the middle of the North Pacific Hilo site, O3 increases at altitudes below 4 km and above 6 km in the troposphere (Fig. S 5(g)), H2O decreases at altitudes below 2 km (Fig. S 5(h)), and temperature decreases throughout the troposphere (Fig. S 5(i)).

We also used ozonesonde data from Hong Kong to test the sensitivity of the regression results when seasonal cycles were excluded. The 7Be data cover 4 winters (2014, 2015, 2016, and 2017), while the ozonesonde data in Hong Kong cover a period of 20 years (2000–2020) and have 20 winters for analysis. Fig. S 6(d), (e), and (f) demonstrate that the profiles of O3 and H2O trends and the increases in the AO over Hong Kong are consistent with the trends of 7Be that vary with AO over Nanning during the analysis period (December 2014–December 2017). Analysis of the 20 years of data shows that O3 increased as the AO increased at altitudes below 7 km (Fig. S 6(a)), H2O decreased as the AO increased at altitudes below 7 km (Fig. S 6(b)), and temperature decreased as the AO increased throughout the troposphere (Fig. S 6(c)). When data only from the winter months (December, January, and February) are considered in the analysis, the results show that increases in O3 occur with increases in the AO at altitudes below 3 km (Fig. S 6(d)), and decreases in H2O occur with increases in the AO at altitudes below 3 km (Fig. S 6(e)). Hence, the results in the lower troposphere show that the O3 and H2O changes with the increases in AO are consistent between all data and winter-only data. Fig. S 6(f) shows that temperatures are slightly increased in the troposphere with increases in the AO. This result is consistent with a longer trend of increases in tropospheric temperatures associated with global warming.

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