Local and Remote Sources of Airborne Suspended Particulate Matter in the Antarctic Region

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Abstract: Quantification of suspended particulate matter (SPM) measurements—together with statistical tools, polar contour maps and backward air mass trajectory analyses—were implemented to better understand the main local and remote sources of contamination in this pristine region. Field campaigns were carried out during the austral summer of 2016–2017 at the “Gabriel de Castilla” Spanish Antarctic Research Station, located on Deception Island (South Shetland Islands, Antarctic). Aerosols were deposited in an air filter through a low-volume sampler and chemically analysed using Inductively Coupled Plasma-Mass Spectrometry (ICP-MS) and Inductively Coupled Plasma-Atomic Emission Spectroscopy (ICP-AES). Elements such as Al, Ca, Fe, K, Mg, Na, P, S, Cu, Pb, Sr, Ti, Zn, Hf, Zr, V, As, Ti, Mn, Sn and Cr were identified. The statistical tools together with their correlations (Sr/Na, Al/Ti, Al/Mn, Al/Sr, Al/Pb, K/P) suggest a potentially significant role of terrestrial inputs for Al, Ti, Mn, Sr and Pb; marine environments for Sr and Na; and biological inputs for K and P. Polar contour graphical maps allowed reproducing wind maps, revealing the biological local distribution of K and P (penguin colony). Additionally, backward trajectory analysis confirmed previous affirmations and atmospheric air masses following the Antarctic circumpolar pattern.

Keywords: Antarctic region; Deception Island; atmospheric aerosols; particulate matter; statistical tools; backward trajectories; polar contour maps

1. Introduction

The Antarctic region is considered to be one of the most virgin, isolated and remote areas globally. The Antarctic is a polar desert and acts as a global thermostat, controlling the global climate system. Nevertheless, although this area is well distanced from continental regions, anthropogenic aerosol pollution from remote and local sources negatively affects the Antarctic environment [1]. Besides, a growing number of tourists visit Antarctica every year. Port Foster, situated in the caldera of Deception Island, is the second most-visited site in this continent, with more than 20,000 visitors in the 2016–2017 austral summer season [2].

Atmospheric aerosols act as climate drivers, since they are involved in the radiation balance of the Earth [3] and the formation of clouds [4]. However, aerosol studies regarding their origin and interactions are still limited and unclear [5]. Aerosols are produced by natural sources (sea, earth erosion, biogenic emissions, volcanoes, etc.) and anthropogenic sources (fossil fuel combustion, mining, agriculture, etc.), negatively affecting the Antarctic’s air quality and ecosystems. Thus, it is important to study them and their impacts on the environment [6].
Atmospheric aerosols have a typical lifetime of one day to two weeks in the troposphere [5]. They transform their size and distribution as they evolve [7], affecting remote places such as the Antarctic [8]. Particulate Matter (PM) transport depends on the wind and meteorological conditions [9,10]. This is important, since the Antarctic region is known for being a trap for aerosols from other places [11].

Air mass back trajectories are precise tools used for investigating the source of the pollutants [12] by estimating the flow patterns of air arriving at the sample collection point. Back trajectory analyses allow for the tracking of the source of aerosols emissions. Long-distance back trajectories are influenced by physical and chemical processes (e.g., deposition and advection), while short-distance trajectories are influenced by emission source zones [13]. However, position inaccuracies of up to 20% of the travel distance can exist [14]. As much as 30% of chemical variability in the troposphere can be associated with transport [15].

Through multivariate analysis, Pérez-Arribas [16] studied a large set of air quality data, aiming to identify elemental correlations and their shared origin. Modelling relationships are important tools to study the relationships between different analytical data, allowing the reproduction of temporal and spatial variations [17,18].

Chemical PM composition has been previously studied in the Antarctic [19]. Nevertheless, despite the importance of PM origin, limited publications using air mass back trajectories exist regarding the Antarctic region [20–22].

The aim of this study is to gain better insight into the potential natural or anthropogenic sources of the measured particulate matter composition and concentration, through air mass back trajectories and polar contour maps. Furthermore, a combination of statistical and graphical approaches is proposed to establish the relationship between the elements and their concentration, in order to reveal the influence of marine environments, human-made activities and penguin colonies on the Antarctic region.

2. Materials and Methods

2.1. Site Description and Sampling

The site description, sampling procedure, and methodology used in the present work, along with the most significant experimental conditions, have been previously described elsewhere [6]. Thus, only the more relevant information to this study is presented here. Atmospheric aerosol particles were collected during the austral summer (from December 2016 to February 2017) at Deception Island (Figure 1), at the Spanish Antarctic Research base “Gabriel de Castilla” (62°58’09” S, 60°42’33” W). A total of 37 samples were collected in circular quartz filter paper of 47 mm diameter (Munktell) by a Derenda LVS 3.1 low volume sampler (2.3 m$^3$/h), and after 24 h, were placed by hand into sterile Petri dishes using sterile tweezers and nitrile gloves. Mass concentration was obtained by gravimetry following the European Norm [23]. Additionally, soil samples were taken from the area of interest. After the gravimetric analysis, Hf, Zr, V, As, Ti, Mn, Cu, Sn, Zn and Pb were determined using Inductively Coupled Plasma-Mass Spectrometry (ICP-MS) and Inductively Coupled Plasma-Atomic Emission Spectroscopy (ICP-AES), as previously described in another publication [6].

2.2. Data Processing

Correlation analysis is a statistical method used to provide information about the relationship between elements that form the air PM by means of a mathematical model. Zhu [24] studied the sources of chemical constituents using multidimensional analysis. The study established correlations between the elements in aerosols. Additionally, a principal component analysis (PCA) was used as an element position simplification tool. Pérez-Arribas [16] adopted PCA to allow a group of correlated data to be transformed into a reduced coordinate system. In this study, Statgraphics Centurion 18, version 18.1.06 of 64-bits (Statpoint Technologies, Warrenton, VA, USA) was used for executing the correlation analysis and the principal component analysis (PCA).
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2.3. Polar Contour Maps

OriginPro 2017, 64-bit (OriginLab Corporation, Northampton, MA, EEUU) was used for studying data relationships through Polar contour maps. These were used as an effective tool for local data visualization. The use of wind direction and speed provided an easy and useful plot of air-pollutants sources. The wind speed is displayed on the x-axis, wind direction on the y-axis of the angle (the radius in degrees), and concentration on the z-axis.

2.4. Air-Mass Back Trajectories

Aiming to distinguish potential remote natural and anthropogenic sources at the sampling location, ten days of air-mass backward trajectories were implemented for an endpoint 600 m above ground level (AGL). This height was selected since the mountainous island has a circular form, and the highest altitude (Mount Pond) is 542 m. In this study, diverse models were used to calculate backward trajectories. Nevertheless, Cabello and Harris [25] pointed out that there are discrepancies between trajectories obtained with different models and different vertical transport methods. The trajectories were calculated with the NOAA HYSPLIT4 (Hybrid Single Particle Lagrangian Integrated Trajectory) model [12,26]. GDAS 1 degree (Global Data Assimilation System) meteorological data from the National Weather Service’s National Centre for Environmental Prediction (NCEP) were chosen as the best appropriate trajectory model over those that did not cover the investigated region and had lower-resolution. GDAS has been widely used to perform backward air trajectories in locations other than the Antarctic region [27–30].

3. Results and Discussion

3.1. Statistical Description of the Data Normality Tests

Particulate matter results show an average of $10 \pm 4 \mu g/m^3$. This value is similar to those found in low population density areas and places such as the Southern Ocean ($13.4 \mu g/m^3$; [31]). However, this result differs from other values found on the Antarctic coast with an average of $1.5 \mu g/m^3$ or $3.4 \mu g/m^3$ [32,33]). The statistical analysis result of total particulate matter (PM10) is shown in Table 1.
The results show a great variability between the sampling days, ranging from 2.9 to 28.2 \( \mu g/m^3 \) with a variation coefficient of 76.4\%. The variability is explained by the singular shape of Deception Island (mountainous horseshoe). This peculiar shape keeps aerosols in the air for an extended period of time.

**Table 1.** Particulate matter (PM) statistical summary together with central tendency, variability, and distribution form.

| Total Particulate Matter (\( \mu g/m^3 \)) | PM10 (\( \mu g/m^3 \)) |
|-------------------------------------------|--------------------------|
| Date                                      |                          |
| 28 December 2016                          | 6.1                      |
| 30 December 2016                          | 28.2                     |
| 01 January 2017                           | 21.7                     |
| 07 January 2017                           | 8.1                      |
| 21 January 2017                           | 12.2                     |
| 22 January 2017                           | 2.9                      |
| 23 January 2017                           | 9.4                      |
| 01 February 2017                          | 4.3                      |
| 08 February 2017                          | 3.8                      |
| 14 February 2017                          | 2.9                      |
| 17 February 2017                          | 4.7                      |
| 23 February 2017                          | 12.1                     |
| 24 February 2017                          | 19.5                     |
| 25 February 2017                          | 7.4                      |
| 26 February 2017                          | 6.0                      |

**PM10 Statistical Overview**

| Number of samples | 15 |
|-------------------|----|
| Average           | 9.95 |
| Standard deviation| 7.60 |
| Coefficient of variation | 76.4\% |
| Minimum           | 2.9 |
| Maximum           | 28.2 |
| Range             | 25.3 |
| Standard Skewness | 2.1299 |
| Standard Kurtosis | 0.8392 |

Both standard skewness and standard kurtosis can be used to determine if the sample has a normal (Gaussian) or different statistical distribution. In general, statistical values outside the range of \(-2\) to \(+2\) indicate significant deviations from normality, which would tend to invalidate many of the statistical procedures that are usually applied to these data.

It can be seen in Table 1 and Figure 2 that the distribution of PM10 data indicates a certain asymmetry, although not well above the critical value of two. Positive asymmetry is usually an indication of a logarithmic-normal or lognormal type of distribution of data. To verify this possibility, the Kolmogorov–Smirnov test was carried out, with the \( P \)-value estimated for a possible lognormal distribution of 0.9983 and 0.6100 estimated for the normal one. Seeing as the two probability values calculated are above the level of usual significance (\( \alpha = 0.05 \)), both statistical distributions are statistically valid. However, as can be seen in Figure 2, the lognormal distribution is better suited to the PM10 data.
obtained. Figure 2 shows a representation of the distribution of number of days (frequency) and PM concentration (μg/m³).

![Histogram of normal and lognormal statistical distribution for the total particulate material obtained during the 2016–2017 field campaign. Frequency indicates the number of days.](image)

Once the general PM statistical behaviour was studied, the same procedure was applied to the different pollutants found in PM10. Specific elements found in the aerosol also showed great variability between different sampling days. However, this variability was not homogeneous for all of them, as shown in Table S1 (Supplementary Materials).

Most of the elements have variation coefficients between 100% and 50%, similar to the PM10 variation coefficient of 76.4%. However, some elements show variation coefficients which are very different and higher than this value. Airborne PM is often affected by different climatic conditions (wind direction, speed, humidity, precipitation, etc.). This implies that, generally, the behaviour of PM composition follows the same trends, but this does not occur for some elements such as; P, K and Sn, with coefficient of variation (C.V.) of 249%, 169.9% and 124.5%, respectively; and for Cu and Fe, with C.V. of 112% and 102%, respectively. This means that these elements may be related to emission sources relatively close to the collection place and possibly very localized, probably due to very intense emission sources. Generally, this situation occurs when the distribution changes from normal or Gaussian statistical behavior, to strong positive asymmetry (lognormal distribution). In the case of K and P, its Kolmogorov–Smirnov test is shown in Table 2 compared to their Gaussian distribution.

| Table 2. P-values obtained for P, K and Cu in PM10 after Kolmogorov–Smirnov test. |
|-----------------------------------|-----------------|-----------------|
|                                   | Lognormal       | Normal          |
| Phosphorus                        | 0.806278        | 0.0123709       |
| Potassium                         | 0.735451        | 0.0267816       |
| Copper                            | 0.288687        | 0.315736        |

In both cases, the P-value for a normal distribution is <0.05, indicating that the contents of K and P in the particulate material are not distributed over the sampling days according to a Gaussian model, and that the lognormal distribution model is much more suitable to describe the behaviour of these major elements. Regarding the trace elements found in the Antarctic aerosol, Cu shows marked positive asymmetry, although in this case, its distribution over the days of sampling can be
explained both by a normal model and by a lognormal model, since the fit tests show $P$-value $> 0.05$ on both models.

3.2. Correlation Analysis

Aiming to explore the chemical/environmental associations among the elements found in the PM10, correlation analysis has been carried out. A statistically significant correlation between elements indicates a common origin of both, so if the source of one of them is identified, it can be presumed that the other comes from the same source of emission. Statistically significant correlations were found between the major constituents in the PM10 [6] and in the case of the elements present at trace levels, 6 significant correlations were found between them [21]. However, it is unknown if some trace elements are correlated to any of the major components. Table S2 (Supplementary Materials) shows the analysis of correlations corresponding to the elements found in the aerosol during the campaign. Those elements that show significant correlation ($P$-value $< 0.05$) are shown in shaded cells.

As Table S2 (Supplementary Materials) shows, there is a very significant correlation between Sr and Na (Pearson $= 0.9873$ and $P$-value $< 0.0001$). Since the main source of Na in the particulate material is marine aerosol, this implies that Sr also has a marine origin. The remaining trace elements do not show a significant correlation with Na, so the marine origin of these elements is discarded. Moreover, Al is a characteristic element of the earth’s crust. Al and Ti have a very significant, high correlation (Pearson $0.9430$; $P$-value $< 0.0001$). Therefore, Ti has a crustal origin as well. Along with its marine origin, Sr has a correlation with Al (Pearson $0.5979$; $P$-value $= 0.0240$). This indicates that some Sr has a terrestrial input. Other trace elements present in the aerosol with possible terrestrial origin are Mn (Pearson $0.8635$; $P$-value $= 0.001$) and Pb (Pearson $0.7557$; $P$-value $0.0028$).

3.3. Multivariate Analysis

Previous correlation analysis has described that there are significant correlations ($P$-value $< 0.05$) among some of the elements found in the PM10. Subsequently, PCA has been carried out, aiming to check if these relationships occur globally. Figure 3 shows the scatter diagrams corresponding to the sampling days. Figure 3a presents the scatter plot when all elements found have been computed as variables; in Figure 3b, the dispersion diagram corresponds to the majority of the elements and Figure 3c corresponds to the components that are at trace levels. Given that the elements were found in different concentration levels, the PCA has been carried out after scaling the data by standard deviation (autoscaling). The green circle indicates that there are two values (samples) close to each other, and therefore there is some similarity between them. However, at the same time, these values are different from the rest. The green and red arrows indicate the same, values that are out of the guideline or general behavior.

As can be seen in Figure 3, the distribution of the scores are quite similar. Most of the points are located at the bottom of the diagram and close to the origin of coordinates. This means that the different pollutants maintain a similar pattern. However, there are exceptions. Figure 3a shows that the sample taken on 1 February 2017 is separated from the rest of the points. This indicates that there is a different pattern compared to the rest of the samples. Similarly, this occurs in Figure 3b,c.

The different patterns show the occurrence of K and P (major elements); and V and as (trace elements). These elements are significantly correlated with each other (see correlation Table S2 in Supplementary Materials). The maximum values of these elements were measured on 1 February 2017, while the rest of the elements gave values close to or below the average. On the other hand, these elements do not show a significant correlation with either Na or Al, so their marine or crustal origin is discarded.
Figure 3. Scatterplot with scores from statistical analysis of the different sampling days by principal component analysis (PCA). (a) represents the whole data set computed as variables. (b) represents the data of the majority elements. (c) represents the data of the trace elements. The green circle, and red and green arrows indicate values that are out of the general behavior.

It can be observed in Figure 3a that for two days (24 February 2017 and 26 February 2017) the elemental composition differs, in relative terms, from the rest of the days. Even though these days appeared close on the distribution diagram, they have different characteristics. One of them corresponds to the presence of majority elements and the other to trace elements. The highest concentration levels were recorded on 24 February 2017 (Figure 3b). The majority elements concentration was recorded to be above the average, except for K and P, on 26 February 2017 (Figure 3c). On this day, a similar situation happened with minority elements, with the exception of V and As (Table S1, Supplementary Materials).

Finally, the point associated to PM collected on 07 January 2017 also deviates from the general pattern. However, this point should be considered anomalous or unrepresentative, since on that day, Fe, Mn, Zn, Sn and Hf were not detected. Furthermore, the rest of the elemental concentration was minimum or largely below the average. Some of these facts could be explained based on the air mass backward trajectories associated with the days prior to the sampling or the sampling meteorological conditions (Figure 4). Generally, air mass backward trajectory analysis showed that PM10 moved
following the Antarctic Circumpolar Pattern through the Southern and Pacific Oceans, while for some of
the cases air masses originated close to the Antarctic Peninsula at the Weddell and Bellingshausen Seas.

- Air mass backward trajectories associated to 01 February 2017 (Figure 4) show previous winds
to this day with origin on the Weddell Sea, passing through the Antarctic continent and the
Bellingshausen Sea. These areas are mostly covered by ice and snow. This explains the low
concentration levels of crustal or marine elements.

- Backward trajectories associated to 24 February 2017 and 26 February 2017 (Figure 4) show
previous days were similar, although on the 26th wind travelled through areas farther north
than on the 24th. In both cases, the wind route passed mainly through ice-free areas, so these
days prevail the presence of Na and the elements correlated with it, such as Mg. In addition, on
26 February 2017 there was moderate rainfall and high relative humidity, which favoured the
deposition of the aerosol.

- On 07 January 2017 (Figure 4), very low levels of all the elements were detected. This cannot be
explained either by the air mass backward trajectories or by the sampling weather conditions.
Therefore, it should be considered to be anomalous.

![Figure 4](image-url)

**Figure 4.** Total 10-day air mass backward trajectories at Gabriel de Castilla Station using GDAS1
(Global data assimilation system) dataset.

### 3.4. Polar Contour Maps

In the statistical study of the pollutant’s distribution, it was found that K and P did not show
Gaussian distribution. However, their distribution was a positive asymmetric lognormal type. This
type of distribution is relatively frequent in the environmental analysis, mainly when there are high
intensity emission sources, relatively close to the sampling site. These sources are usually located
somewhere very localized.

Since the transport of pollutants in the air is carried out by the wind, polar contour maps were
used to relate the concentration of these pollutants in the air with wind direction and speed. On the
other hand, according to the correlation study mentioned above, K and P have an important and
significant correlation between them (Pearson 0.9821; $P$-value < 0.0001). Alternatively, they do not
present a significant correlation with either Na or Al. This indicates that both elements have the same
non-crustal, marine origin.
If we look at the polar contour maps associated to these two elements (Figure 5), the emission focus of both maps is in the 210° direction (south-west of the sampling site). This implies a natural and biological origin of these elements, since the Punta de la Descubierta penguin colony is located in that direction. This colony is one of the largest penguin colonies in Deception Island, so the presence of these elements in the PM is related to the excrement (guano) in the area.

![Figure 5. (a) Polar contour maps in relation to wind direction and speed for phosphorus (P) and (b) potassium (K) at the Gabriel de Castilla Spanish Research Station (Deception Island).](image)

4. Conclusions

By using statistical methods combined with polar contour Maps and air mass back trajectories, the different sources of PM collected during the austral summer campaign in 2016–2017 (2.5-month term) at the Spanish Antarctic Research Base “Gabriel de Castilla” located on Deception Island have been determined.

Correlation analysis between major and minor elements showed the potential source of the airborne PM. Sr/Na correlation indicated the influence of marine aerosols, whereas Al/Ti correlation revealed the influence of local crustal sources. Al/Mn, Al/Sr and Al/Pb correlations indicated terrestrial sources. Furthermore, K/P correlation was revealed to have non-crustal/marine origins, since no correlation with Na or Al was found. Multivariate analysis proved the importance of establishing guidelines in the behaviour of PM10 and its composition throughout the sampling period. Score distribution showed a similar pattern for most of the elements, except for K and P. The origin of the K/P correlation was validated by PCA and polar contour maps to be biological (penguin scats from the penguin colony).

Air mass back trajectories were used to confirm the elemental source. These trajectories revealed that both crustal and marine inputs occurred following different pathways and were influenced by the Antarctic Circumpolar pattern.

This work revealed the importance of an improved understanding of the potential origin and behaviour of PM in the Antarctic, through the use of statistical tools, air mass back trajectories and polar contour maps. Consistent aerosol tracking in the Antarctic region is crucial, since atmospheric aerosols play a significant role in the Earth’s climate and ecosystems. Further work will be undertaken on Deception Island as well as at other Antarctic stations.

Supplementary Materials: The following are available online at http://www.mdpi.com/2073-4433/11/4/373/s1.

Table S1: Statistical summary of major (a; µg/m³) and minor elements (b; ng/m³). Those elements that show significant deviation from the Gaussian model (statistical values outside the range −2 to +2) are in red., Table S2: Statistics of correlations between different elements. For each element the first, second and third row correspond to Pearson correlation coefficient, sample size and P-Value, respectively. Those elements that show significant correlation (P-value <0.05) are shown in shaded cells.

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**References**

1. Turner, J.; Barrand, N.E.; Bracegirdle, T.J.; Convey, P.; Hodgson, D.A.; Jarvis, M.; Jenkins, A.; Marshall, G.; Meredith, M.P.; Roscoe, H.; et al. Antarctic climate change and the environment: An update. *Polar Rec.* 2014, 50, 237–259. [CrossRef]

2. IAATO. *Report on IAATO Operator Use of Antarctic Peninsula Landing Sites and Atcm Visitor Site Guidelines, 2016-17 Season.* Ip 164; International Association of Antarctica Tour Operators (IAATO): South Kingston, RI, USA, 2017; Available online: https://iaato.org/es/past-iaato-information-papers (accessed on 28 April 2017).

3. Nielsen, I.E.; Skov, H.; Massling, A.; Eriksson, A.C.; Dall’Osto, M.; Junninen, H.; Sarnela, N.; Lange, R.; Collier, S.; Zhang, Q.; et al. Biogenic and anthropogenic sources of aerosols at the High Arctic site Villum Research Station. *Atmos. Chem. Phys.* 2019, 19, 10239–10256. [CrossRef]

4. Tomasi, C.; Lanconelli, C.; Mazzola, M.; Lupi, A. Aerosol and Climate Change: Direct and Indirect Aerosol Effects on Climate. In *Atmospheric Aerosols*; Wiley-VCH Verlag GmbH & Co. KGaA: Weinheim, Germany, 2016; pp. 437–551.

5. IPCC. *Climate Change 2013: The Physical Science Basis. Contribution of Working Group i to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change*; Cambridge University Press: Cambridge, UK; New York, NY, USA, 2013.

6. Cáceres, J.O.; Sanz-Mangas, D.; Manzoor, S.; Pérez-Arribas, L.V.; Anzano, J. Quantification of particulate matter, tracking the origin and relationship between elements for the environmental monitoring of the Antarctic region. *Sci. Total Environ.* 2019, 665, 125–132. [CrossRef] [PubMed]

7. Mahowald, N.; Albani, S.; Kok, J.F.; Engelstaeder, S.; Scanza, R.; Ward, D.S.; Flanner, M.G. The size distribution of desert dust aerosols and its impact on the Earth system. *Aeolian Res.* 2014, 15, 53–71. [CrossRef]

8. Hu, Q.-H.; Xie, Z.-Q.; Wang, X.-M.; Kang, H.; Zhang, F. Levoglucosan indicates high levels of biomass burning aerosols over oceans from the Arctic to Antarctic. *Sci. Rep.* 2013, 3, 3119. [CrossRef]

9. Radojevic, M.; Bashkin, V.N. *Practical Environmental Analysis*; RSC: Cambridge, UK, 2006.

10. Reeve, R.N. *Introduction to Environmental Analysis*; John Wiley & Sons: Chichester, UK, 2002.

11. Bargagli, R. Atmospheric chemistry of mercury in Antarctica and the role of cryptogams to assess deposition patterns in coastal ice-free areas. *Chemosphere* 2016, 163, 202–208. [CrossRef]

12. Stein, A.F.; Draxler, R.R.; Rolph, G.D.; Stunder, B.J.B.; Cohen, M.D.; Ngan, F. NOAA’s HYSPLIT Atmospheric Transport and Dispersion Modeling System. *Bull. Am. Meteorol. Soc.* 2015, 96, 2059–2077. [CrossRef]

13. Fleming, Z.L.; Monks, P.S.; Manning, A.J. Review: Untangling the influence of air-mass history in interpreting observed atmospheric composition. *Atmos. Res.* 2012, 104, 1–39. [CrossRef]

14. Hondula, D.M.; Sitka, L.; Davis, R.E.; Knight, D.B.; Gawtry, S.D.; Deaton, M.L.; Lee, T.R.; Normile, C.P.; Stenger, P.J. A back-trajectory and air mass climatology for the Northern Shenandoah Valley, USA. *Int. J. Climatol.* 2010, 30, 569–581. [CrossRef]

15. Moody, J.; Galusky, J.; Galloway, J. The use of atmospheric transport pattern recognition techniques in understanding variation in precipitation chemistry. In *Atmospheric Deposition (Proceedings of the Baltimore Symposium, May 1989)*; IAHS Publ. No. 179; IAHS: Wallingford, UK, 1989.

16. Pérez-Arribas, L.V.; León-González, M.E.; Rosales-Conrado, N. Learning Principal Component Analysis by Using Data from Air Quality Networks. *J. Chem. Educ.* 2017, 94, 458–464. [CrossRef]
17. Cristofanelli, P.; Putero, D.; Bonasoni, P.; Busetto, M.; Calzolari, F.; Camporeale, G.; Grigioni, P.; Lupi, A.; Petkov, B.; Traversi, R.; et al. Analysis of multi-year near-surface ozone observations at the WMO/GAW “Concordia” station (75°06′S, 123°20′E, 3280 m a.s.l. – Antarctica). *Atmos. Environ.* 2018, 177, 54–63. [CrossRef]

18. Mihalikova, M.; Kirkwood, S. Tropopause fold occurrence rates over the Antarctic station Troll (72°S, 2.5°E). *Ann. Geophys.* 2013, 31, 591–598. [CrossRef]

19. Osipov, E.Y.; Osipova, O.P.; Khodzher, T.V. Recent variability of atmospheric circulation patterns inferred from East Antarctica glaciochemical records. *Geochemistry* 2019, 125544. [CrossRef]

20. Mishra, V.K.; Kim, K.-H.; Hong, S.; Lee, K. Aerosol composition and its sources at the King Sejong Station, Antarctic peninsula. *Atmos. Environ.* 2004, 38, 4069–4084. [CrossRef]

21. Marina-Montes, C.; Pérez-Arribas, L.V.; Escudero, M.; Anzano, J.; Cáceres, J.O. Heavy metal transport and evolution of atmospheric aerosols in the Antarctic region. *Sci. Total Environ.* 2020, 721, 137702. [CrossRef]

22. Szumirska, D.; Czapiewski, S.; Szopińska, M.; Polkowska, Z. Analysis of air mass back trajectories with present and historical volcanic activity and anthropogenic compounds to infer pollution sources in the South Shetland Islands (Antarctica). *Bull. Geogr. Phys. Geogr. Ser.* 2018, 15, 111–137.

23. European Committee for Standardization (CEN). *UNE EN 12341:2015. Ambient Air. Standard Gravimetric Measurement Method for the Determination of the PM10 and PM2.5 Mass Concentration of Suspended Particulate Matter*; European Committee for Standardization: Brussel, Belgium, 2015.

24. Zhu, G.; Guo, Q.; Xiao, H.; Chen, T.; Yang, J. Multivariate statistical and lead isotopic analyses approach to identify heavy metal sources in topsoil from the industrial zone of Beijing Capital Iron and Steel Factory. *Environ. Sci. Pollut. Res.* 2017, 24, 14877–14888. [CrossRef]

25. Cabello, M.; Orza, J.A.G.; Gallano, V.; Ruiz, G. Influence of meteorological input data on backtrajectory cluster analysis – a seven-year study for southeastern Spain. *Adv. Sci. Res.* 2008, 2, 65–70. [CrossRef]

26. Rolph, G.; Stein, A.; Stunder, B. Real-time Environmental Applications and Display System: READY. *Environ. Model. Softw.* 2018, 95, 210–228. [CrossRef]

27. Jorba, O.; Pérez, C.; Roca­denbosch, F.; Balda­sano, J. Cluster Analysis of 4-Day Back Trajectories Arriving in the Barcelona Area, Spain, from 1997 to 2002. *J. Appl. Meteorol.* 2004, 43, 887–901. [CrossRef]

28. Moore, B.J.; Neiman, P.J.; Ralph, F.M.; Barthold, F.E. Physical Processes Associated with Heavy Flooding Rainfall in Nashville, Tennessee, and Vicinity during 1–2 May 2010: The Role of an Atmospheric River and Mesoscale Convective Systems. *Mon. Weather Rev.* 2012, 140, 358–378. [CrossRef]

29. Su, L.; Yuan, Z.; Fung, J.C.H.; Lau, A.K.H. A comparison of HYPLIT backward trajectories generated from two GDAS datasets. *Sci. Total Environ.* 2015, 506–507, 527–537. [CrossRef] [PubMed]

30. Hong, S.-b.; Yoon, Y.; Becagli, S.; Gim, Y.; Chambers, S.D.; Park, K.-T.; Park, S.-J.; Traversi, R.; Severi, M.; Vitale, V.; et al. Seasonality of aerosol chemical composition at King Sejong Station (Antarctic Peninsula) in 2013. *Atmos. Environ.* 2019, 117185. [CrossRef]

31. Budhavant, K.; Safai, P.D.; Rao, P.S.P. Sources and elemental composition of summer aerosols in the Larsemann Hills (Antarctica). *Environ. Sci. Pollut. Res.* 2015, 22, 2041–2050. [CrossRef]

32. Budhavant, K.B.; Rao, P.S.P.; Safai, P.D. Size distribution and chemical composition of summer aerosols over Southern Ocean and the Antarctic region. *J. Atmos. Chem.* 2017, 74, 491–503. [CrossRef]

33. Mazzera, D.; Lowenthal, D.; Chow, J.; Watson, J.; Grubišić, V. PM10 measurements at McMurdo Station, Antarctica. *Atmos. Environ.* 2001, 35, 1891–1902. [CrossRef]