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Mass Concentration, Chemical Composition, and Source Characteristics of PM$_{2.5}$ in a Plateau Slope City in Southwest China

Jianwu Shi $^{1}$, Yinchuan Feng $^{1}$, Liang Ren $^{1}$, Xiuqing Lu $^{2}$, Yaoqian Zhong $^{1}$, Xinyu Han $^{1,2,*}$ and Ping Ning $^{1}$

1 Faculty of Environmental Science and Engineering, Kunming University of Science and Technology, Kunming 650500, China; Shijianwu@kust.edu.cn (J.S.); renliang@stu.kust.edu.cn (L.R.); zhong@kust.edu.cn (Y.Z.); ningping58@sina.com (P.N.)
2 Architectural Engineering Institute, Kunming University of Science and Technology, Kunming 650500, China; Lxq08258@126.com

* Correspondence: 20110020@kust.edu.cn; Tel.: +86-15912128009

Abstract: In order to investigate the seasonal variations in the chemical characteristics of PM$_{2.5}$ at the plateau slope of a mountain city in southwest China, 178 PM$_{2.5}$ filters (89 quartz and 89 Teflon samples for PM$_{2.5}$) were collected to sample the urban air of Wenshan in spring and autumn 2016 at three sites. The mass concentrations, water-soluble inorganic ions, organic and inorganic carbon concentrations, and inorganic elements constituting PM$_{2.5}$ were determined, principal component analysis was used to identify potential sources of PM$_{2.5}$, and the backward trajectory model was used to calculate the contribution of the long-distance transmission of air particles to the Wenshan area. The average concentration of PM$_{2.5}$ in spring and autumn was 44.85 ± 10.99 µg/m$^3$. Secondary inorganic aerosols contributed 21.82% and 16.50% of the total PM$_{2.5}$ in spring and autumn, respectively. The daily mean value of OC/EC indicated that the measured SOC content was generated by the photochemical processes active during the sampling days. However, elements from anthropogenic sources (Ti, Si, Ca, Fe, Al, K, Mg, Na, Sb, Zn, P, Pb, Mn, As and Cu) accounted for 99.38% and 99.24% of the total inorganic elements in spring and autumn, respectively. Finally, source apportionment showed that SIA, dust, industry, biomass burning, motor vehicle emissions and copper smelting emissions constituted the major components in Wenshan. This study is the first to investigate the chemical characteristics and sources of PM$_{2.5}$ in Wenshan, and it provides effective support for local governments formulating air pollution control policies.

Keywords: PM$_{2.5}$; PCA; backward trajectories; chemical composition; source

1. Introduction

In recent decades, with rapid economic development, industrialization and urbanization in China, the number of motor vehicles and the total energy consumption have increased, and atmospheric particulate matter (PM) has become one of the most significant air contaminants [1–3]. PM, particularly PM$_{2.5}$ (aerodynamic diameter ≤ 2.5 mm), can exist in the atmosphere for a long time, which is conducive to its long-distance transport through the atmosphere and deposition towards remote areas. During long-range transport, PM$_{2.5}$ carries abundant anthropogenic pollutants and has a serious impact on the global and regional climate, the visibility and composition of the atmosphere, the global biogeochemical cycle and the activation of cloud condensation nuclei (CCN) [4–6].

PM$_{2.5}$ has been widely studied in recent years in China due to its potential impacts on air quality and human health. Water-soluble inorganic ions (WSIs), organic carbon (OC) particles, inorganic carbon (EC) particles, and inorganic elements (IEs), as the main chemical components of PM$_{2.5}$, have been extensively studied in China [7–9]. WSIs are dominated by secondary inorganic aerosols (SIA), including NH$_4^+$, NO$_3^-$, and SO$_4^{2−}$. 

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OC is composed of thousands of organic compounds and contains many toxic substances. Heavy metals are an important part of the inorganic elements comprising PM$_{2.5}$, such as Fe, Zn, Cu and Pb [10–12].

Whether worldwide or only in China, it is essential to reduce PM$_{2.5}$ concentrations to control their sources. The key point in formulating policies for the government to control PM$_{2.5}$ pollution is the result of source apportionment and reliable source quantification [13,14]. In fact, PM$_{2.5}$ is usually sourced by the emission of pollutants, and its classification is very complex, including its anthropogenic and natural sources and gas and particle phases [15,16]. In addition, PM$_{2.5}$ forms secondary pollutants from primary emissions through photochemical reactions after being released from pollution sources, making it difficult to quantify its impacts [17–19]. The contribution levels of different sources in the air can be quantitatively estimated by using the receptor model. Generally, researchers have identified the possible sources of PM$_{2.5}$ as traffic and industrial, coal combustion, biomass burning and secondary inorganic aerosol sources [20,21].

In recent years, most studies have generally focused on the Jing-Jin-Ji region and its coastal areas with severely degraded atmospheric environments in China [22–24]. Only a few researchers have investigated PM$_{2.5}$ pollution in Yunnan Province, which is a remote southwestern region. More research has been conducted in areas such as Kunming and Yuxi [25–27]. Despite the economic backwardness of the remote southwestern mountains, there are still cases of PM$_{2.5}$ exceeding the standard every year [28–30]. Therefore, we should pay more attention to these areas to improve their air quality.

Wenshan, a developing industrialized city in southwest China, has a high degree of air pollution, mainly resulting from the presence of PM$_{2.5}$ in the atmosphere. Furthermore, Wenshan is located in the basin valley on the plateau, and the urban area is surrounded by high mountains, which aggravate particulate pollution. Wenshan is chosen as the study area to conduct PM$_{2.5}$ sampling during the spring and autumn seasons at three monitoring sites. The concentrations of PM$_{2.5}$, WSIs, OC, EC and IEs are analyzed and discussed in the current study. Principal component analysis (PCA) is used for PM$_{2.5}$ source apportionment to analyze the pollution sources. Potential major contributors were identified on the basis of PCA and local environmental background information. The details of the pollution characteristics and the results of PM$_{2.5}$ source apportionment in this study can provide the local government with reasonable and effective measures to slow down atmospheric pollution with PM$_{2.5}$.

2. Methods

2.1. Sampling Site and Sample Collection

Wenshan is a developing industrialized city with half a million inhabitants in an urban area of 75 km$^2$. With a longitude of 103$^\circ$43′–104$^\circ$27′ E and a latitude of 23$^\circ$06′–23$^\circ$44′ N, Wenshan lies in southwest China (Figure 1), which is the transition zone of the Yunnan-Guizhou Plateau and Vietnam Basin.

Wenshan lies in a small basin valley on the plateau and is surrounded by mountains on three sides. The terrain inclines from northwest to southeast, and the mountain range runs almost from north to south. Therefore, a corridor topography consisting of high points on both sides and low points in the middle is formed. The relative altitude difference is 1751.2 m, with the highest altitude of 2991.2 m in Bozhu Mountain and the lowest of 1240 m in the urban area. Wenshan is dominated by a subtropical monsoon climate, which is characterized by a long spring and autumn, no bitter cold in winter, no brutal heat in summer and delightful weather in all seasons. Airflow near the ground can only enter the urban area from the southeast. The cold air in Siberia from the north is obstructed by mountains, and the monsoon of the Beibu Gulf and the Bay of Bengal traveling from the southeast flows right into the urban area, which forms comfortable temperatures, low wind speeds and high ultraviolet (UV) light conditions throughout the year. Strong ultraviolet light is conducive to the formation of photochemical atmospheric effects, and the conditions of low pressure and low oxygen can lead to the incomplete combustion of fuel and can
increase the resulting pollution emissions. The conditions of low wind speed (<3 m/s) and the large diurnal temperature variation readily form an inversion layer, hindering the diffusion of pollutants.

![Location of the sampling site in Wenshan, China.](image)

Measurement campaigns of PM$_{2.5}$ sampling at three sites at the Convenience Service Center of Wenshan (CSCW), Water Authority of Wenshan (WAW) and Environmental Monitoring Station of Wenshan (EMSW) in Wenshan city (Figure 1), were carried out in two periods in 2016, i.e., 19 April to 3 May and 12 October to 26 October. Daily measurements of 22 ± 1 h with an intelligent mid-volume atmospheric particulate sampler (TH-150F, Tianhong, China) were conducted at a 100 L/min sample flow. Teflon filters (China, 90 mm) were used to analyze OC particles, EC components and WSIs. A total of 178 PM$_{2.5}$ samples and 12 blank samples were collected. After sampling, the filters were individually placed into plastic boxes and put into a freezer at −20°C until transport and subsequent analysis.

2.2. Chemical Analysis and Quality Control

2.2.1. WSII Analysis

The anion (i.e., F$^-$, Cl$^-$, SO$_4^{2-}$ and NO$_3^-$) and cation (i.e., K$^+$, Ca$^{2+}$, Mg$^{2+}$ and NH$_4^+$) concentrations were measured by ion chromatography (DX-600, Dionex, USA). This system was outfitted with a separation column (Dionex AS-14A for anions and CS-12A for cations) and a guard column (Dionex AG14A for anions and CG12A for cations). A gradient weak base eluent (3.5 mmol/L Na$_2$CO$_3$; 1 mmol/L NaHCO$_3$) was used for anion detection, while a weak acid eluent (18 mmol/L methanesulfonic acid) was used for cation detection. The measurement error of each ion in a standard solution is within 10%, and the average relative standard deviations of anions and cations are 3.0% and 4.0%, respectively. For quality assurance, two blank spaces were detected in each batch of samples, and the test was carried out at 10%. At least six standard solution concentrations needed to be mixed for each ion component.

2.2.2. Elements Analysis

Li, Be, P, Cr, Mn, Bi, Co, Ni, Cu, Sr, Mo, Cs, Cd, Tl, Pb, Th, Sc, V, As, Rb, Y, Zr, Sn, Sb, La, Ce, U, Sm and W were analyzed by inductively coupled plasma-mass spectrometry (ICP-MS). Mg, Ca, K, Fe, Al, Ba, Zn, Na and Si were analyzed by inductively coupled plasma-atomic emission spectrometry (ICP-AES).
2.2.3. OC and EC Analysis

The concentrations of OC and EC were analyzed by DRI Model 2001 OC/EC Analyser, which was developed by the American Desert Institute (DRI). The main testing principle of this method is as follows: the sample is heated and converted into CO$_2$ under different temperature gradients and gas environments. CO$_2$ is reduced to CH$_4$ by catalyzing MnO$_2$ and is detected by using a flame ion detector (FID). Then, using a 633 nm helium/neon laser to detect the anti-light intensity of filter paper to detect the production of organic pyrolysis carbon (OPC), eight different carbon components (OC1, OC2, OC3, OC4, OPC, EC1, EC2, and EC2) were obtained. IMPROVE (Interagency Monitoring of Protected Visual Environments) defines OC as OC1+OC2+OC3+OC4+OCPyro and EC as EC1+EC2+EC3-OCPyro. The detection limits were 0.82 (OC), 0.19 (EC) and 0.93 (TC) µg/cm$^2$, and the measuring range was 0.2~750 µg/cm$^2$.

2.2.4. Principal Component Analysis (PCA) Modeling

PCA is an important multivariate statistical tool that can reduce the dimensionality of large datasets and extract the number of principal components needed to explain all the variance of such datasets, which is much less than the original number of variables [31,32]. PCA extracts new variables by the correlation between all variables, which contain most of the information about the data, called principal components. Each variable has the same significance, and each topic has the same weight. The first component extracted explains the maximum amount of data variance. The maximum amount of remaining data variance will be further explained by successive components [33–35]. This process sets up the orthogonal distribution of components to each other, and the result of the regression adjustment of factors is simple and stable, regardless of how large a dataset is and how many variables are included in the study [36].

2.2.5. HYSPLIT4 Model

The HYSPLIT4 model is a professional model jointly developed by the National Oceanic and Atmospheric Administration (NOAA) Air Resources Laboratory (ARL) and the Australian Bureau of Meteorology (BOM) in the past 20 years to calculate and analyze the transport and diffusion trajectories of atmospheric pollutants. The model has a relatively complete transport, diffusion and sedimentation model that can handle multiple meteorological element input fields, multiple physical processes, and different types of pollutant emission sources. It is widely used in the study of the transmission and diffusion of multiple pollutants in various regions. In this study, the independent version of the backward trajectory model was used, the auxiliary software package was used (GUI; Ghostscript; ImageMagick), and the meteorological data were obtained through NCEP (National Centers for Environmental Prediction) and GDAS (Global Data Assimilation System).

2.2.6. Quality Control

The quartz filters were baked at 450 °C for 5 h in a muffle furnace before sampling to identify the possible presence of organics. All filters were placed in a clean room (temperature of 25 °C ± 5; relative humidity of 50 ± 5%) for 48 h and weighed by a high-precision electronic balance (EX125ZH) with an accuracy of 10 mg before and after sampling. Each filter was weighed twice, with the difference between the two results not exceeding 0.2 mg for quartz filters and 0.02 mg for Teflon filters, to guarantee the precision of the weighting results. All filters were stored in a freezer at −20 °C before analysis to prevent the loss of volatile components.

In the sample analysis process, the instrument was calibrated with standard gas before and after the sample analysis. Then, one sample was randomly selected from every 10 samples for parallel analysis, and the standard sample was measured twice a week. The recovery rate of the standard sample was 98%~102%. Finally, the system blank of the instrument and the blank of the laboratory system were measured every week. The results
showed no contamination during sample handling and collection, as assured by the quality assurance and control (QC/QA) procedures.

3. Results and Analyses

3.1. Concentration Characteristics of PM$_{2.5}$

During the two sampling campaigns, the mass concentrations (from Telfon filters) of PM$_{2.5}$ ranged from 29.11 $\mu$g/m$^3$ to 62.03 $\mu$g/m$^3$ in spring and from 24.46 $\mu$g/m$^3$ to 60.08 $\mu$g/m$^3$ in autumn (Figure 2). The overall concentration of PM$_{2.5}$ is higher in spring than that in autumn. The concentration on individual days in autumn is higher than that in spring, which may be related to changes in meteorological conditions. The daily concentration levels of PM$_{2.5}$ were all within Chinese National Ambient Air Quality Standards II (75 $\mu$g/m$^3$). In addition, the total average concentration of PM$_{2.5}$ in spring and autumn (44.85 $\pm$ 10.99 $\mu$g/m$^3$) was higher than Standard II (35.00 $\mu$g/m$^3$) (GB3095-2012), and was 3.0 times higher than the annual standard concentration in the USA (15 $\mu$g/m$^3$). These values are lower than those of developed cities in the plains of China, such as Beijing, Tianjing and Shanghai [37,38]. Furthermore, the concentration of PM$_{2.5}$ in Wenshan was lower than that in some plateau cities, such as Guiyang and Kunming [39]. In our previous research, we found that the dust emission volume of Wenshan was 1164 t, accounting for 52.9% of the total emissions. PM$_{2.5}$ pollution may be associated with city construction, and the floor space of buildings under construction was 5.48 $\times$ 10$^6$ m$^2$ during 2016 in Wenshan. The mean concentrations of PM$_{2.5}$ decreased from spring (48.00 $\pm$ 11.01 $\mu$g/m$^3$) to autumn (41.64 $\pm$ 10.10 $\mu$g/m$^3$). There were 10 and 5 days in spring and autumn, respectively, that exceeded the total average concentration, which means that were 62.5% of the days in spring exceeded the total average concentration, and it was twice that in autumn. This seasonal mean concentration variation is attributed to the primary influence of meteorological characteristics and source emissions.

Figure 2. The daily mass concentration of PM$_{2.5}$ during sampling.

3.2. Chemical Composition Characteristic of PM$_{2.5}$

3.2.1. WSIIs Levels

The concentrations of WSIIs were 11.75 $\pm$ 3.25 and 12.50 $\pm$ 3.40 $\mu$g/m$^3$ in spring and autumn, respectively. In PM$_{2.5}$, the concentrations of K$^+$, NH$_4^+$, Ca$^{2+}$, Mg$^{2+}$, Cl$^-$, F$^-$, NO$_3^-$ and SO$_4^{2-}$ were 0.67, 2.54, 1.28, 0.07, 0.24, 0.16, 1.25 and 5.53 $\mu$g/m$^3$ in spring and 0.59, 3.17, 0.78, 0.07, 0.38, 0.22, 0.77 and 6.53 $\mu$g/m$^3$ in autumn, respectively. The annual concentration of WSIIs was 12.15 $\mu$g/m$^3$ and occupied 26.91% of the total PM$_{2.5}$. This result indicated that WSIIs were one of the main components of PM$_{2.5}$. The mass concentrations of sulfate occupied 49.67% of the total WSIIs, followed by NH$_4^+$ (23.50%), Ca$^{2+}$ (8.60%), NO$_3^-$ (8.37%), K$^+$ (5.21%), Cl$^-$ (2.52%), F$^-$ (1.56%), and Mg$^{2+}$ (0.57%). The dominant compounds were secondary inorganic aerosols (SIAs, including NO$_3^-$, SO$_4^{2-}$,
and NH$_4^+$, with concentrations accounting for more than 80% of the total WSII mass of PM$_{2.5}$. The concentrations of WSIs in Wenshan are shown in Table 1 and compared to other typical cities, such as Kunming [30] and Guiyang [40] on the plateau and Beijing [41] and Nanjing [42] on the plain. Compared to other Chinese studies, most of the ionic species identified in research are found to be on the lower side. Compared with Kunming and Guiyang, the concentration of SO$_4^{2-}$ was lowest, which is consistent with the lagged industrial development of Wenshan. These results show that the WSII concentrations at Wenshan were impacted more by local pollution sources (e.g., biomass burning, agricultural dust, construction dust, etc.) [43].

**Table 1.** Mean concentrations of WSIs sampled in Wenshan in 2016 compared with data from other sites (µg/m$^3$).

| Ion   | Wenshan         | Kunming [30] | Guiyang [40] | Beijing [41] | Nanjing [42] |
|-------|-----------------|--------------|--------------|--------------|--------------|
| K$^+$ | 0.63 ± 0.24     | 0.77         | 0.41         | 1.90         | 1.2          |
| Ca$^{2+}$ | 1.04 ± 0.5   | 2.83         | 1.77         | 3.92         | 0.7          |
| Mg$^{2+}$ | 0.07 ± 0.02   | 0.30         | 0.20         | 0.76         | 0.2          |
| NH$_4^+$ | 2.81 ± 1.16  | 0.52         | 4.29         | 12.47        | 4.5          |
| Cl$^-$ | 0.31 ± 0.19     | 0.72         | 1.30         | 6.76         | 1.9          |
| F$^-$  | 0.19 ± 0.07     | 0.54         | 0.03         | -            | 0.2          |
| SO$_4^{2-}$ | 5.98 ± 2.07  | 9.72         | 17.43        | 21.60        | 5.1          |
| NO$_3^-$ | 1.00 ± 0.40   | 0.51         | 1.34         | 20.552       | 9.1          |

SIAs were the dominant ions in the PM$_{2.5}$ component in both autumn and spring. The spring and autumn concentrations of SIAs follow the order SO$_4^{2-}$ > NH$_4^+$ > NO$_3^-$ (Figure 3). One of the reasons is that industrial production leads to the incomplete combustion of fossil fuels, which increases the emission of the gaseous precursor SO$_2$ [44,45]. Moreover, the geographical structure of urban areas is not favorable to pollutant diffusion in the atmosphere. Another reason is the high conversion rate of SO$_2$ to PM$_{2.5}$, which may be due to the relatively high humidity in autumn [46,47]. In addition, NH$_4^+$ was the most dominant cation in PM$_{2.5}$ in the two seasons, and the emission of NH$_4^+$ originated from the nitrogen fertilizers used in agriculture [48,49]. The observed NO$_3^-$ levels were related to the synthetic action of various influencing factors, i.e., precursor NO$_x$ emissions, complex photochemical and heterogeneous reactions and gas-aerosol equilibrium [50,51].

**Figure 3.** Seasonal variations in SIAs and their ratios in PM$_{2.5}$ in Wenshan.

To discuss the relative importance of mobile and stationary sources of SO$_2$ and NO$_x$, the mass concentration ratio of NO$_3^-$/SO$_4^{2-}$ was used as an indicator [52]. The seasonal variation in NO$_3^-$/SO$_4^{2-}$ in PM$_{2.5}$ ranged from 0.16 to 0.32 and from 0.09 to 0.18 in spring and autumn, respectively, with an annual mean of 0.18 ± 0.07, which was lower than the values measured in Shanghai (0.43) [53], Qingdao (0.35) [54] and Taiwan (0.20) [55].
Therefore, with the increasing number of motor vehicles, the contribution of mobile sources is more important than before.

Ion balance calculations are frequently used to investigate the acid base balance of ions in PM$_{2.5}$. The correlation of CE and AE and the variation in CE/AE in the two seasons were calculated. According to the electroneutrality of solutions, AE must be equal to CE [56]. The correlation coefficient between CE and AE for spring ($R^2 = 0.92$) was higher than that for autumn ($R^2 = 0.85$), showing that cations and anions maintained better equilibrium during neutralization in spring. The average CE/AE ratios for autumn (1.72) were higher than those for spring (1.54), which indicates the basic nature of aerosols in which PM$_{2.5}$ is alkaline in the two seasons in Wenshan [57,58].

3.2.2. IEs Levels

The concentrations of IEs in PM$_{2.5}$ in the two seasons are shown in Figure 4. The concentrations of IEs in PM$_{2.5}$ were 4.82 µg/m$^3$ and 4.10 µg/m$^3$ in spring and autumn, respectively. Fifteen main elements, Ti, Si, Ca, Fe, Al, K, Mg, Na, Sb, Zn, P, Pb, Mn, As and Cu, account for 99.38% and 99.24% of the total inorganic elements in spring and autumn, respectively. These fifteen elements play a key role in the estimation of emission sources and are associated with human activity (such as industrial processes, residential activities, and traffic patterns).

![Figure 4. Mean concentrations of inorganic elements in PM$_{2.5}$ sampled at Wenshan.](image)

In the PM$_{2.5}$ samples, the relatively high concentrations of elements are in the order of Ti>Si>Ca>Al>Fe>K>Mg>Na>Sb>Zn>P>Pb>Mn>As>Cu (spring) and Ti>Si>Ca>Fe>Al>K>Mn>Mg>Na>Zn>Sb>Pb>P>As>Cu (autumn). The fifteen main IE concentrations accounted for 14.88% of the total PM$_{2.5}$ in spring and 14.89% in autumn. The concentrations of the identified elements of soil dust (Ti, Si, Al, Ca, Mg) were 5.24 µg/m$^3$ in spring and 4.29 µg/m$^3$ in autumn, which showed that surface dust was the main source of PM$_{2.5}$. The concentrations of the industrial discharge elements (As, Zn, and Mn) were 0.10 µg/m$^3$ in spring and 0.38 µg/m$^3$ in autumn. The concentration of Pb was 0.03 µg/m$^3$ in spring and 0.06 µg/m$^3$ in autumn, and it may be due to motor vehicle exhaust emissions. The concentrations of K accounted for 1.35% of PM$_{2.5}$ in spring and 0.99% in autumn, which may be due to biomass burning [59–61].

3.2.3. OC and EC Levels

The mean concentrations of OC were 12.03 ± 2.24 µg/m$^3$ and 9.32 ± 2.13 µg/m$^3$ in spring and autumn, respectively (Figure 5). The mean EC concentrations were 3.66 ± 0.47 µg/m$^3$ and 2.88 ± 0.61 µg/m$^3$ in spring and autumn, respectively. Wenshan is located in the basin valley on the plateau, with wind speeds that are too low to be conducive to pollutant spreading during the two seasons. During the two sampling campaigns, the daily mean value of OC/EC was 2.64–4.17 in spring and 2.74–3.65 in autumn, all of which exceeded 2.0, which indicated that Wenshan experienced secondary organic carbon (SOC) pollution in both seasons. Moreover, OC and EC in Wenshan had a better correlation in autumn ($R = 0.86$) than in spring ($R = 0.69$), which showed that the measured OC and
EC were derived from similar sources during autumn and from complex sources during spring. The possible reason is that the spring is affected by the long-distance transmission of biomass combustion sources in Southeast Asia due to climatic conditions.

Since there is no simple and direct calculation method for SOC, this study estimates the content of SOC considering the lowest value of the OC/EC ratio in the two seasons [62–64]. The principle of this method is the use of the lowest value of OC/EC rather that of the OC/EC of the primary pollutant in every season. SOC = TOC, EC × (OC/EC)min, and TOC and EC are the concentrations of OC and EC in PM2.5, respectively [65,66]. The average values of SOC in PM2.5 are 2.36 ± 1.00 μg/m3 in spring and 0.99% in autumn, respectively. Wenshan is located in the basin valley on the plateau, with wind speeds that are too low to be conducive to pollutant spreading during the two seasons. During the two sampling campaigns, the radiation and temperature in autumn were higher than those in spring, which represented more favorable photochemical conditions for the formation of SOC in autumn. However, there was more rain in autumn in Wenshan, which could limit the formation of SOC. The order of OC, SOC and SOC/EC was the same pattern as spring > autumn, which meant that the impact of radiation and temperature was less than that of rainfall, and rainfall was the main influencing factor and had a greater impact on SOC [67,68].

3.3. Source Apportionment of PM2.5
3.3.1. Principal Component Analysis (PCA)

The sample quantity is crucial for good PCA. The PCA of the study was performed considering the chemical constituents of 90 PM2.5 samples. The outliers (those beyond 2SD) were removed, and the dataset was normalized prior to the operation [69]. When the value of KMO is close to 1, it indicates that there is a strong correlation between these variables (KMO indicates the amount of variance shared among the items designed to measure a latent variable when compared to that shared with the error), and these variables are more suitable for PCA [70]. In this study, the SPSS software package (IBM, version 24.0) was used to conduct PCA research on substances in PM2.5 to obtain the emission characteristics.
of its pollution sources. For this study, the species of Li, Na, K, Mn, Cu, Zn, As, Pb, Al, Mg, Ca, Fe, Ba, Si, Ti, \( \text{NH}_4^+ \), \( \text{NO}_3^- \), \( \text{SO}_4^{2-} \), OC and EC had strong correlations in the two seasons, and the PCA results are listed in Table 2.

**Table 2.** Matrix of loading factors of PCA in spring and autumn.

| Element | Spring | Autumn |
|---------|--------|--------|
| Li      | 0.52   | 0.86   |
| Na      | 0.80   | 0.52   |
| K       | 0.93   | 0.77   |
| Mn      | 0.71   | 0.88   |
| Cu      | 0.94   | 0.76   |
| Zn      | 0.62   | 0.91   |
| Pb      | 0.64   | 0.55   |
| Al      | 0.56   | 0.82   |
| Mg      | 0.93   | 0.78   |
| Ca      | 0.91   | 0.76   |
| Ba      | 0.90   | 0.77   |
| Fe      | 0.87   | 0.74   |
| Si      | 0.70   | 0.65   |

From Table 2, we know that PCA resolved six components explaining 80.5% of the variance in spring.

**Factor 1:** The first factor contributes 18.15% to the total factor contributions and is characterized by a high concentration of SIAs, which indicates that Wenshan was greatly affected by secondary inorganic aerosol pollution in spring. SIAs are mainly generated by the photochemical reactions of precursor gases (SO\(_2\), NH\(_3\), and NO\(_x\)), which are emitted from specific identified sources of human activity (coal combustion, vehicle exhaust emission, and biomass burning). Therefore, the strict control of precursor gases is conducive to reducing \( \text{PM}_{2.5} \) levels.

**Factor 2:** The second factor contributed 16.28% of the total \( \text{PM}_{2.5} \), and mostly originated from natural sources, such as the lifting of dust or mechanical abrasion processes, which was identified by high concentrations of Al, Fe, Si and Ti, indicating the leading contribution of dust [71]. Si and Ti are the key tracers of soil dust caused by winds. The extra Ba is emitted from brake linings and tire tread wear. These results can be explained as a consequence of dust persisting in the atmosphere longer because of calm and low-speed winds.

**Factor 3:** The third factor resolved 14.15%, and represents the factor contribution from industrial emissions. The elements are related to the industrial metal smelting process and represent anthropogenic emissions from various industries near the sampling site [72].

**Factor 4:** This source provided 11.55% of \( \text{PM}_{2.5} \). OC and EC are considered to be tracers of motor vehicle emissions, and EC is an indicator of primary emissions of OC [73]. The presence of K also deserved our attention, directly indicating emissions from biomass burning.

**Factor 5:** Cu and Na were apportioned to this factor, which suggests that the effect of this factor was manifold, such as copper smelting and sea salt [74]. The contribution of this factor towards \( \text{PM}_{2.5} \) was 10.83%, as revealed by PCA. In addition, Na might travel long
Distances from the Indian Ocean, and Cu could have come from the nearby industrial area in Honghe Prefecture.

**Factor 6:** This factor is construction cement dust, which is represented by high concentrations of Ca and Mg [75]. This finding indicates that construction and demolition activities were prevalent in the urban areas in Wenshan during the sampling period, without effective measures for dust control. More precise and effective policies are needed for the local government to improve PM$_{2.5}$ pollution.

In addition, PCA resolved five components explaining 78.7% of the variance in autumn. Different from spring, Factor 1 represents biomass burning and industry sources, contributing 29.40% of PM$_{2.5}$. Factor 2 includes secondary inorganic aerosols and motor vehicle exhaust emissions, which resolved 18.73% of the factor contribution. Factor 3 represents metal smelting, with remarkable representative Al, Fe and Mg features, which can be attributed to the smelting production activities around the site. Factor 4 and Factor 5 are soil dust and construction dust, which resolved 9.30% and 8.91% of the factor contributions, respectively.

### 3.3.2. The Long-Range Transport

To better understand the transport of airborne particles from distant sources, the 72 h backward trajectories starting at a height of 100 m at the sampling site were calculated using the Hybrid Single Particle Lagrangian Integrated Trajectory 4.0 (HYSPLIT4) model with a 12 h period (meteorological data from the Global Data Assimilation System (GDAS)). The back trajectories were classified into three clusters using TrajStat in this study.

In spring, the trajectories were grouped into three clusters (Figure 6). Cluster 1 (blue line), from the southwestern direction, was associated with slower and lower air mass trajectories and accounted for 57%. The other two trajectory clusters (green line and red line) came from the north and southwestern directions, accounting for 24% and 19%, respectively. Cluster 1 came from central Myanmar and passed through during spring sampling in northern Vietnam and the Honghe Prefecture in Yunnan Province, China, which explains the effect of Factor 3. At the same time, Cluster 2 (green line) came from the industrial region in Chongqing, which explained the source of biomass burning in Factor 1 and the industrial impact in Factor 3. Figure 6b shows that wind mainly originated from the south during the sampling period, which prevented the diffusion and great accumulation of NO$_x$ and SO$_2$. Then, they formed secondary pollution through photochemical reaction transformation, which conforms to the SIA pollution in Factor 1. The higher wind speeds were also consistent with the contribution of Factor 2.

In autumn, the trajectories were grouped into three clusters from the southeastern direction (Figure 7). Cluster 1 (red line) came from Guangxi Province and passed through the industrial region in Baise, which explains the presence of industrial elements in Factor 1. The other two trajectory clusters (blue line and green line) accounted for 33% and 8%, respectively. Figure 7b also shows that the wind mainly originated from the south during the sampling period and resulted in the impossibility of the diffusion and dilution of pollutants, which was also the reason for the SIA pollution in Factor 2.
the south during the sampling period, which prevented the diffusion and great accumulation of NO\textsubscript{X} and SO\textsubscript{2}. Then, they formed secondary pollution through photochemical reaction transformation, which conforms to the SIA pollution in Factor 1. The higher wind speeds were also consistent with the contribution of Factor 2.

Figure 6. (a) Mean 72 h backward trajectories of each trajectory cluster during spring and the percentage of allocation to each cluster. (b) Wind roses of Wenshan during spring sampling.

In autumn, the trajectories were grouped into three clusters from the southeastern direction (Figure 7). Cluster 1 (red line) came from Guangxi Province and passed through the industrial region in Baise, which explains the presence of industrial elements in Factor 1. The other two trajectory clusters (blue line and green line) accounted for 33% and 8%, respectively. Figure 7b also shows that the wind mainly originated from the south during the sampling period and resulted in the impossibility of the diffusion and dilution of pollutants, which was also the reason for the SIA pollution in Factor 2.

Figure 7. (a) The mean 72 h backward trajectories of each trajectory cluster during autumn and the percentage of allocation to each cluster. (b) Wind roses of Wenshan during autumn sampling.

4. Conclusions

In this study, PM\textsubscript{2.5} samples were collected in Wenshan, and their mass concentration, chemical composition and source apportionment characteristics were analyzed in spring and autumn. The mean concentrations of PM\textsubscript{2.5} were 48.00 ± 11.01 μg/m\textsuperscript{3} and 41.64 ± 10.10 μg/m\textsuperscript{3} in spring (sampled on 19 April–3 May) and autumn (sampled on 12 October–26 October). The annual mean concentration of PM\textsubscript{2.5} at the three sites was 44.85 ± 10.99 μg/m\textsuperscript{3}, which was lower than that in Standard II (75.00 μg/m\textsuperscript{3}) and higher
than that in Standard II (35.00 µg/m³). This means that the air quality in Wenshan is better than that in most cities in China.

WSIIs and OC were the main components of PM$_{2.5}$, accounting for 26.91% and 23.80% of PM$_{2.5}$, respectively. SIAs were the major contributors to WSIIs, due to the incomplete combustion of fossil fuels and the slathering of nitrogen fertilizers in agriculture. Wenshan was greatly affected by secondary inorganic aerosol pollution in the two seasons, which contributed 21.82% and 16.50% to the total factor contributions in spring and autumn, respectively. The ratio of NO$_3^−$/SO$_4^{2−}$ implied that the contribution of mobile sources was not significantly different from that of other developed areas. The daily mean value of OC/EC was 2.64–4.17 in spring and 2.74–3.65 in autumn, which indicates that the SOC was generated by the photochemical process during the sampling days in Wenshan. Moreover, the OC and EC concentrations in Wenshan had a better correlation in autumn (R = 0.86) than in spring (R = 0.69), which shows that OC and EC were derived from similar sources during autumn and from complex sources during spring. However, elements from anthropogenic sources (Ti, Si, Ca, Fe, Al, K, Mg, Na, Sb, Zn, P, Pb, Mn, As and Cu) accounted for 99.38% and 99.24% of the total inorganic element concentration in spring and autumn, respectively.

Source apportionment showed that SIAs (18.15%), the lifting of dust or mechanical abrasion processes (16.28%), industrial sources (14.15%), motor vehicle emissions (11.55%), copper smelting and sea salt pathways (10.83%), and construction cement dust emissions (9.56%) were the main pollution sources in PM$_{2.5}$ in spring. Furthermore, source apportionment showed that biomass burning and industry (29.40%), SIAs and motor vehicle exhaust (18.73%), metal smelting (12.33%), soil dust (9.30%) and construction dust (8.91%) emissions were the main pollution sources of PM$_{2.5}$ in autumn. Different source contributions were found in spring and autumn. According to the research results, the pollution prevention and control suggestions are as follows: (1) Exert related effective management for artificial sources, such as industry and construction sites, to accelerate industrial transformation and upgrading. (2) Adopt emission control measures, such as motor vehicle restrictions and the promotion of new energy transportation methods.

The results of cluster analysis indicate that the long-range transport of air pollutants has a profound effect on local air quality in Wenshan. Wenshan is mainly affected by long-distance atmospheric transmission from the southwest and the northeast in spring and autumn, respectively.

In this paper, chemical composition and source characteristics of PM$_{2.5}$ in a plateau slope city were first studied, and the main sources of PM$_{2.5}$ in Wenshan City are resolved. The results can provide scientific data to support PM$_{2.5}$ pollution control in local and similar cities.

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Nomenclature

PM$_{2.5}$: Particulate matter with aerodynamic equivalent diameter less than or equal to 2.5 microns in ambient air.

NOx: Refers to the sum of NO and NO$_2$.

CCN: Cloud condensation nuclei.

WSIs: Water-soluble inorganic ions.

OC: Organic carbon.

EC: Inorganic carbon.

IEs: Inorganic elements.

PCA: Principal component analysis.

UV: Ultraviolet.

CSCW: Convenience Service Center of Wenshan.

WAW: Water Authority of Wenshan.

EMSW: Environmental Monitoring Station of Wenshan.

ICP-AES: Inductively coupled plasma-atomic emission spectrometry.

DRI: Desert Institute.

FID: Flame ion detector.

OPC: Organic pyrolysis carbon.

IMPROVE: Interagency Monitoring of Protected Visual Environments.

SIAs: Secondary inorganic aerosols (including NO$_3^-$, SO$_4^{2-}$ and NH$_4^+$).

CE/AE: Cation/Anion concentration ratio.

NCEP: National Centers for Environmental Prediction.

GDAS: Global Data Assimilation System.

QC/QA: Quality control and Quality assurance.

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