Submicron Aerosol and Black Carbon in the Troposphere of Southwestern Siberia (1997–2018)

Mikhail Panchenko 1, Elena Yausheva 1,*, Dmitry Chernov 1, Valerii Kozlov 1, Valery Makarov 2, Svetlana Popova 2 and Vladimir Shmargunov 1

1 V.E. Zuev Institute of Atmospheric Optics, Siberian Branch, Russian Academy of Sciences, 1, Ak. Zuev square, 634021 Tomsk, Russia; pmv@iao.ru (M.P.); chernov@iao.ru (D.C.); vkozlov@iao.ru (V.K.); vpsh@iao.ru (V.S.)
2 V.V. Voevodsky Institute of Chemical Kinetics and Combustion, Siberian Branch, Russian Academy of Sciences, Institutskaya 3, 630090 Novosibirsk, Russia; makarov@kinetics.nsc.ru (V.M.); popova@kinetics.nsc.ru (S.P.)

Abstract: Based on the multiyear measurements in the surface atmospheric layer (from five stations) and regular flights of aircraft laboratory over the background region of Southwestern Siberia, the compositions of mass concentrations of submicron aerosol and absorbing substances (soot and black carbon) are analyzed. The annual average concentrations of submicron aerosol and black carbon were found to be maximal in 1997, 2012, and 2016, when the largest numbers of wildfires occurred across the entire territory of Siberia. No significant, unidirectional trend of interannual variations in the concentration of submicron particles was observed, while the concentration of absorbing substance reliably decreased by 1.5% each year. To estimate the effect of urban pollutants, mass concentrations of aerosol and absorbing substance in the surface layer at the Aerosol Station (in the suburban region of Tomsk) were compared to those at the Fonovaya Observatory (in the background region). It was shown that the largest contribution of anthropogenic sources in the suburban region was observed in the winter season, while minimal difference was observed in the warm period of the year. The seasonal behavior of the concentrations of elemental carbon at three stations in Novosibirsk Oblast almost completely matched the dynamics of the variations in the black carbon concentration in the atmosphere of Tomsk Oblast. Data of aircraft sensing in the troposphere of the background region of Southwestern Siberia (2000–2018) were used to determine the average values of the vertical distribution of the submicron aerosol and black carbon concentrations in the altitude range of 0.5–7 km for each season. It was found that at altitudes of 0.5–7 km, there were no unidirectional trends in submicron aerosol; however, there was an increase of black carbon concentration at all altitudes with a positive trend of 5.3 ± 2.2% per year at an altitude of 1.5 km, significant at a p-value = 0.05.

Keywords: aerosol; black carbon; elemental carbon; aircraft measurements; trends; wildfires

1. Introduction

Under conditions of rapid changes in the climate system and intensifying anthropogenic effects, the problem of correctly accounting for and predicting the aerosol effect on environmental state, human health, and the formation of Earth’s radiation budget still urgently needs a large amount of data on the properties of atmospheric particles from instrumental observations [1].

The main difficulties when determining the aerosol effect are not only associated with the strong spatiotemporal variations in the concentration of particles and their microstructural and physicochemical characteristics, but also with a complex set an intricate complex of multidirectional processes, in which aerosol plays a leading role.
Anthropogenic aerosol has a negative effect on nature and human health, especially in big megacities; therefore, solutions to this serious problem have attracted the attention of the entire global community (see e.g., [2–9]).

For the state of radiation balance of the troposphere, it is possible to single out the following main processes of both direct radiative forcing and indirect aerosol effects. The direct radiative forcing of particles is directly related to their optical characteristics, where the most important aspect is the relationship between the scattering and absorbing properties. Scattering has a cooling effect, while absorption has a warming effect [10,11]. In the case of indirect aerosol effects, the main roles are played by the concentration, chemical composition, and particle size distribution. Functioning as the condensation nuclei in cloud formation (cooling factor), aerosol particles determine the number of cloud droplets and the microstructure, and, correspondingly, their optical properties and lifetimes in the atmosphere [12,13].

The indirect deferred, cumulative aerosol effect is another indirect aerosol effect, the study of which recently attracted the attention of the scientific community regarding the extremely rapid changes observed in the climate system of the Arctic [14–20]. This effect is caused by the deposition of absorbing particles on the underlying surface, the accumulation of which leads to a reduction in the surface albedo. The effect of this process is most urgent for the northern territories, where the arrival of absorbing substances in ice and snow cover substantially increases not only the absorption of radiation, but also ice and snow melt.

To solve climatically significant problems, the present-day scientific community has implemented a great number of large-scale international projects worldwide, for which comprehensive research into aerosol properties at different spatial scales has already been carried out for a few decades (e.g., ARM, AERONET, and GAW [21–23]).

Obtaining reliable information on the properties of scattering and absorbing particles in the troposphere at a regional scale is one of the most urgent and complex problems [24–26]. In this case, studying particles in a submicron size range comes to the forefront. First, submicron particles mainly determine the scattering and absorption of incoming shortwave radiation. This size range also comprises the main mass of strongly absorbing soot particles (black carbon (BC)), the content of which usually does not exceed 10% of the total aerosol mass concentration. However, it is these particles that play an important role in both direct radiative forcing and in indirect aerosol effects [27], and they are the second most important global warming agent after greenhouse gases. Second, submicron particles have the largest lifetimes in the troposphere, and are transported long distances away. Hence, their microphysical composition is gradually transformed and depends on the territory of their origin and the trajectories of air mass motion; therefore, their effect on the properties of atmospheric air is manifested on considerable spatial scales far away from a specific measurement site.

In this work, we discuss the results from multiyear studies of submicron aerosol and black carbon concentrations in the troposphere of Southwestern Siberia. Based only on a separate series of observations, it is unrealistic to identify regularities of interannual, seasonal, and diurnal variations in aerosol characteristics ("aerosol weather" [28]) under the influence of a complex combination of synoptic and meteorological processes. Weather is defined as a physical state of the atmosphere at a certain instant or in a finite period of time, determined by a set of meteorological quantities and atmospheric phenomena. Aerosol is an integral part of the atmosphere and, as such, heavily determines its optical state and should generally be considered as an element of weather.

Satellite observations and measurements of the aerosol optical depth with the application of sun photometers are known to be feasible only under the conditions of clear-sky and low-cloud weather. These data provide information for a limited set of atmospheric situations (as applied to conditions of Southwestern Siberia, there are only 15–20% cases, observed usually in anticyclones and in Arctic air masses). Thus, this problem can only be solved using continuous, multiyear observations in a monitoring mode. Here, we carried out a regional scale analysis to determine the aerosol states correctly using observations
at five ground-based stations in the Tomsk and Novosibirsk Oblasts. Data from regular monthly flights of an aircraft laboratory over a background region in Novosibirsk Oblast were employed to estimate the interannual variations in the troposphere of the region.

2. Method and Instruments

Here, we will make just a few important methodic clarifications. The formation of the aerosol composition in the troposphere is known to depend on many processes, namely: on the geographic location of the origin of air mass and, correspondingly, on the strength of the sources and the sinks on the trajectory of the air mass motion to the observation region, i.e., processes with a relatively long impact period (crudely 3–10 days). To exclude the effect of fast-flowing processes of change in the relative air humidity with a pronounced diurnal behavior on the transformation of dry matter of submicron particles, the flow of the analyzed air, taken from the atmosphere before being admitted to the instruments, was artificially dried through heating at 20$^\circ$C from ambient.

Taking into account the main problem being addressed in this work, among the entire set of characteristics observed, we employed measurements of the angular scattering coefficient of the dry aerosol matter, $\mu$ (45$^\circ$), and mass concentrations of black carbon and elemental carbon (EC) for analysis.

Photoelectric aerosol nephelometers (PhANs) measured the angular scattering coefficient of the dry matter of the submicron particles $\mu_d$ (45$^\circ$) (Mm$^{-1}$ sr$^{-1}$) at an angle of 45$^\circ$ with a sensitivity level of molecular scattering of ~1 Mm$^{-1}$ sr$^{-1}$. The instruments were calibrated against molecular scattering by pure air and CO$_2$ gas. The angular scattering nephelometer of PhAN-A provided measurements using a lamp as the source of radiation at an effective wavelength of 0.51 µm. The modified PhAN-M nephelometer had a light diode as a source of radiation at a wavelength of 0.53 µm [28,29].

The volume scattering coefficient was estimated using the empirical formula $\sigma_d = 7.3 \cdot \mu_d$ (45$^\circ$) (Mm$^{-1}$). For the convenience of our researchers studying the aerosol processes through non-optical means (such as through gravimetry), the mass concentration of the submicron aerosol was estimated according to a one-parameter aerosol optical model $M_A$ ($\mu g/m^3$) = 2.4 $\cdot$ $\mu_d$ (45$^\circ$) (for a particle density of 1.5 g/cm$^3$) [29].

The concentration BC$_{eq}$ was measured using a multiwavelength differential aethalometer (MDA) designed at the Institute of Atmospheric Optics, Siberian Branch, Russian Academy of Sciences (IAO SB RAS), with a sensitivity of ~20 ng/m$^3$ when pumping 20 L of air through the device (Table 1). The concentration of BC$_{eq}$ was measured using a multiwavelength differential aethalometer (MDA) designed at IAO SB RAS, with a sensitivity of ~20 ng/m$^3$ when pumping 20 L of air through the device. It is noted that the aethalometer measured the concentration of the absorbing substance in the aerosol in units of equivalent mass concentration BC$_{eq}$ ($\mu g/m^3$), because the instrument was calibrated with respect to black carbon particles in order to determine the mass on the basis of recording the aerosol absorption.

Table 1. Locations and characteristics of monitoring stations.

| Station Name       | Site Type       | Location       | Period        | Method                  | Characteristic | Time Resolution |
|--------------------|-----------------|----------------|---------------|-------------------------|----------------|-----------------|
| Aerosol Station    | Suburban        | 56.48$^\circ$ N, 85.05$^\circ$ E | 1997–2018    | Nephelometry, aethalometry | $M_A$, BC$_{eq}$ | hourly         |
| Fonovaya Observatory| Background      | 56.40$^\circ$ N, 84.07$^\circ$ E | 2014–2018    | Nephelometry, aethalometry | $M_A$, BC$_{eq}$ | hourly         |
| Novosibirsk        | Urban center    | 55.01$^\circ$ N, 82.57$^\circ$ E | 2000–2016    | Thermal method          | EC             | 1 day (1 month in season) |
| Klutchi            | Suburban        | 54.51$^\circ$ N, 83.16$^\circ$ E | 1999–2017    | Thermal method          | EC             | 1 day (1 month in season) |
| Zavyalovo          | Background      | 54.26$^\circ$ N, 82.18$^\circ$ E | 2001–2016    | Thermal method          | EC             | 1 day (1 month in summer) |
| Troposphere (aircraft) | Background     | Novosibirsk Oblast | 2000–2018    | Nephelometry and aethalometry | $M_A$ and BC$_{eq}$ | once a month (0.5–7 km) |
To determine the elemental carbon (EC) in the composition of the surface aerosol, samples were collected on fiberglass (Whatman®) filters 47 mm in diameter at a rate of 30 l/min over 24 h. The method of reaction gas chromatography was used to determine the diurnally average concentrations of the EC. This method made it possible to determine the EC content through its high-temperature separation from organic carbon in an inert-oxidizing environment. Each component was oxidized to give CO$_2$, converted to CH$_4$, and was recorded using a flame-ionization detector. Thus, when sample was heated (700 °C) in an inert atmosphere, the organic compounds evaporated and the organic carbon was determined; meanwhile, the elemental carbon was quantified during combustion in the oxidizing atmosphere [30].

3. Measurement Sites

3.1. Aerosol Station

The Aerosol Station at the Institute of Atmospheric Optics, Siberian Branch, Russian Academy of Sciences (AS IAO SB RAS) is located in Akademgorodok, a suburban region of Tomsk (Figure 1); at this site, the measurements of this set of aerosol characteristics has been carried out in monitoring mode since 1997 [31]. The angular scattering coefficient of the submicron particles $\mu$ (45°) (Mm$^{-1}$ sr$^{-1}$) was determined using a PhAN-A nephelometer. The concentration BC$_{eq}$ was measured using a multiwavelength differential aethalometer (MDA).

![Figure 1. Geographic positions of the measurement sites and flight paths of the aircraft laboratory (color lines). Google Earth (Image Landsat/Copernicus).](image)

3.2. Fonovaya Observatory

The Fonovaya Observatory is located in a forested zone on the coast of the River Ob, 4 km from the nearest small village, Kireevsk. Tomsk is 60 km east of the measurement site. Given the predominant southwesterly and southerly winds in the region, the main wind direction from Tomsk does not have an effect on this terrain; therefore, this territory experi-
ences much less anthropogenic load and, as such, can be considered as a conventionally background region.

Since November 2013, the PhAN-M nephelometer measurements of the angular scattering coefficient have been carried out at the observatory every hour in a monitoring mode, and the BC_{eq} concentration has been measured using MAAP 5012 aethalometer (Thermo Scientific Fisher Inc., Franklin, TN, USA).

3.3. Novosibirsk

The measurement site is located at the center of Novosibirsk. Its population of approximately 1.5 million people occupies an area of 502 km². The main local sources of anthropogenic aerosol in this region are motor vehicles (approximately 400,000) and thermal power plants. The distance between Tomsk and Novosibirsk is 205 km. Atmospheric air samples for aerosol analysis were taken from 2000 to 2016 every year, in winter (20 January–19 February), spring (20 April–19 May), summer (20 June–19 July), and autumn (20 September–19 October).

3.4. Klutchi Village

The site is located in Klutchi village, 30 km southeast of Novosibirsk, and is considered rural terrain. Its population of approximately 100 people occupies an area of 11 km². It is thought that the composition of absorbing substance in aerosol in this area is formed as a result of natural sources slightly contaminated by pollutants of an anthropogenic origin. Here, from 1999 to 2017, samples were collected to determine the EC, at the time intervals as in Novosibirsk.

3.5. Zavyalovo Village

The Zavyalovo sampling site is located in a forested zone, 90 km southeast of Novosibirsk and 15 km from the nearest small Zavyalovo village. In this terrain, there are no local sources of anthropogenic pollution from the atmosphere. Aerosol samples were taken here from 2001 to 2016 in order to determine the concentrations of EC in the summer season, primarily because of the lack of a suitable infrastructure for collecting aerosol samples during the cold seasons.

3.6. Aircraft Laboratory

Mass concentrations of submicron aerosol and BC_{eq} were measured since 2000 on board aircraft laboratory, in the framework of complex studies of atmospheric composition.

The flights of the aircraft laboratory (Optik-E AN-30 from 1999 to 2008, and Optik TU-134 since 2011) were carried out once a month, all year round, during daytime under clear-sky conditions.

The Optik-E AN-30 aircraft laboratory was chosen as a research platform because of several advantages it has. It is a high-wing, two turboprop airplane, which has one more turbojet engine in the right nacelle. It can fly in a wide range of altitudes from 100 to 7000 m with speeds of 70–125 m/s. Another advantage is that this aircraft was specifically designed for airborne mapping; therefore, it has five optical windows in the bottom, which can be used for remote sensing [32].

Because the AN-30 aircraft ended its operational life in 2011, the question about choosing another aircraft for placing the equipment and continuing research arose. The closest in terms of technical characteristics was the TU-134 aircraft—a short-range passenger aircraft with two turbojet engines, which was the basis of the new Optic TU-134 aircraft laboratory.

The studied route passes over a pine forest along the right-hand coast of the Ob Reservoir, near Zavyalovo Village, in the southern region of Novosibirsk oblast (Figure 1). This terrain was chosen as the proving ground for background measurements, because cities and industrial activities, being sources of anthropogenic aerosols, are far away from this territory.
Sensing was carried out according to the following pattern: when approaching the region of measurements, the aircraft climbed to altitude of 7 km, then, circling over a specified area, it gradually descended (on a horizontal flight leg, signals were recorded for 8 min altitudes of 7.0, 5.5, 4.0, 3.0, 2.0, 1.5, 1.0, and 0.5 km). Flights were carried out once a month, each lasting, on the average, for approximately 3 h. A modified nephelometer PhAN-M was used on board the aircraft laboratory to measure the aerosol scattering coefficient, and the MDA was used to measure the \( BC_{eq} \) concentration. In total, 142 flights of the aircraft laboratory were carried out over the period of 2000–2018, which were analyzed here.

4. Results and Discussion

Before discussing the results from the analysis of the multiyear measurements, which were required to develop prognostic models, the following points must be noted.

In order for the multiyear time series of surface measurements to be used to determine the characteristic properties of the tropospheric aerosol on a specifically regional scale, the effect of forest fires should be identified first.

Wildfires are an integral component of regional “aerosol weather” in Siberia, the territory of which is mostly covered by forest massifs [13,33–38]. This factor ambiguously affects the atmospheric radiation regime, because large amounts of both non-absorbing particles and black carbon are emitted into the atmosphere during biomass burning. Smoke from remote wildfires in the surface atmospheric layer are known to be a cooling factor (direct radiative forcing) [33,34]. It is noteworthy that the absorption of radiation in the upper layers of the smoke mass leads to its warming and lifting of air up to altitudes of cloud formation, which contains a greater concentration of particles, serving mostly as condensation nuclei [19,35,36]. Accordingly, formation of thick low-level clouds intensifies the cooling of lower atmospheric layers (indirect aerosol effects).

The large volume of heated smoke mass and clouds is transported large distances and can penetrate as far as remote regions of the Arctic. Under the influence of processes of dry and wet deposition, absorbing aerosol particles, after being supplied to the underlying surface, decrease the surface reflectivity, leading to longer-term and more serious consequences, and serving as the warming factor (indirect deferred, cumulative aerosol effect) [39–42]. To estimate the contribution of wildfires to changes in the concentrations of submicron aerosol and black carbon, an analysis was carried out for the entire ensemble of data and for the dataset, in which the events with smoke haze present in the atmosphere over the observation sites were excluded (without fires). The situations with the presence of smokes were selected on the basis of estimates of the parameter \( P = \frac{BC_{eq}}{M_A} \), i.e., the \( BC_{eq} \) fraction of submicron aerosol, and through an analysis of the satellite information. Based on the multiyear measurements at the Aerosol Station, the stable regularity revealed a lower \( BC_{eq} \) fraction in forest fire smoke (1 to 5%) compared with that in the background aerosol, i.e., they were weakly absorbing [29]. The conventionally called \( P \)-effect was present in the decrease of the \( P \) values in conditions of smoke haze relative to their values in natural conditions. This effect is caused by the dominating contribution of the pyrolysis of wooden combustible materials to the formation of properties of smoke mist particles. The value of the \( BC_{eq} \) fraction of the dry matter of submicron aerosol \( P \) was determined as the ratio of the mass concentrations of \( BC_{eq} \) and aerosol, a with the mean error of about 20–25%. The effect of smoke from forest fires was established using a fire map from [43], composed on the basis of satellite information and an analysis of the meteorological situation.

The second factor, which influences the properties of tropospheric aerosol, analyzed on the basis of a multiyear time series of surface measurements, was the anthropogenic sources, located near the measurement sites. This problem was solved quite easily by comparing synchronous measurements at urban and suburban sites against data obtained by the stations in background regions.
4.1. Seasonal Variations
4.1.1. Surface Measurements

Effect of Forest Fires

Figure 2 presents the multiyear seasonal behavior of mass concentrations of submicron aerosol and black carbon at the Aerosol Station (suburban region). Red boxes denote the total datasets, comprising measurements loaded by the effect of remote forest fires. Green and black boxes show the average mass concentrations of submicron aerosol and black carbon, processed to remove the events loaded by smoke from forest fires. After the datasets of MA and BC$_{eq}$ were processed to eliminate the effect of forest fires, their maximal values in the surface atmospheric layer were observed in a cold period of the year, and minimal values were noted in summer. The reasons for the annual behavior will be discussed below.

As seen, the annual behavior of the measured characteristics is affected appreciably by smoke from remote forest fires, occurring from April to October, in the Siberian region.

It should be noted that during the fire-dangerous period, smoke from remote wildfires was observed most often in May and July. The effect of smoke plumes from forest fires over the entire period of observations at the Aerosol Station led to increased average concentrations of aerosol and BC in this month, e.g., by factors of 2 and 1.5, respectively, at the Aerosol Station. Figure 3 presents the interannual variations in the identified fire situations in these months. From 1997 to 2018, there were fewer days in May with smoke haze present, probably owing to the prohibition issued on combustion of the previous year’s vegetation. On the contrary, in July, there were more days with smoke from forest fires. Analogous results were also obtained in the literature [44], based on which, over the period of 2002–2018, a negative trend of aerosol optical depth (AOD) in spring was identified, as well as increasing tendencies for its summertime values. Atmospheric AOD measurements were carried out all year round in the vicinity of Tomsk using a solar photometers [22,45].

Figure 2. Multiyear seasonal behavior of mass concentrations of (a) submicron aerosol and (b) black carbon (BC)$_{eq}$ at the Aerosol Station (suburban region of Tomsk) in 1997–2018. Red boxes indicate the total dataset. Green and black boxes are for datasets without accounting for the contribution of situations with wildfires. The lower and upper sides of the box show the 25 and 75 percentile concentrations, respectively; the horizontal bars represent the median; and the closed squares give the mean. The whiskers represent the 5th and 95th percentile concentrations.
The longest-term and severest smoke situations were observed in 1997, 2012, and 2016, when Southwestern Siberia was covered by smoke plumes from extensive forest fires for a long time. The maximal daily mass concentrations of submicron aerosol and black carbon were observed at the Aerosol Station on 27 July 2012, when the diurnal mass concentration of submicron aerosol was 1213 μg/m³ and the BC_{eq} concentration was 27 μg/m³. The situation was the same in the EC concentration measurements at three sites in Novosibirsk Oblast, with the strongest smoke pollution of the atmosphere observed in July 2012. The measurements of aerosol characteristics in this period recorded a substantially increased fraction of submicron aerosol and organic carbon (OC) [46]. The EC concentrations in the period of maximal smoke pollution of the atmosphere reached 26 μg/m³ in Novosibirsk, 11 μg/m³ in Klutchi, and 6 μg/m³ in Zavyalovo.

Estimates of the Effect of Urban Aerosol Sources

To estimate the effect of urban aerosol sources, a comparative analysis of the values at ground-based sites was carried out without accounting for situations with the effect of smoke from remote forest fires. It should be noted that the average multiyear seasonal behaviors of M_A and BC_{eq} at the suburban station (Aerosol Station) and in the background region (Fonovaya Observatory) were close in shape, being maximal in winter and minimal in the warm period of the year (Figure 4).

However, as expected, the concentrations of aerosol and BC_{eq} at the sites, subject to a stronger anthropogenic load, were much higher, and the amplitude of their seasonal behavior was more pronounced.

For the average annual behavior at the Aerosol Station (Tomsk) over the entire period of 1997–2018, the concentration of submicron aerosol M_A was maximal (32 ± 9.9 μg/m³) in February and minimal (13.7 ± 2.9 μg/m³) in September (Figure 2a, Table 2). In the atmosphere over the Fonovaya Observatory over the period of 2014–2018 (period of synchronous measurements at two stations), the average aerosol concentration was also maximal (21.6 ± 3.4 μg/m³) in February and minimal (7.3 ± 1.7 μg/m³) in May (Figure 4a). In this same period, at the Aerosol Station, the maximal average concentration, recorded in February, increased to 41.4 ± 14.4 μg/m³, and the minimal concentration, revealed in May, was 10.1 ± 1.5 μg/m³. Thus, the largest contribution of anthropogenic sources in the suburban region over 2014–2018 was observed in winter season: in January and February, the aerosol concentration was, on the average, 20 μg/m³ (a factor of 1.9) larger.
than for the background conditions. A minimal difference was revealed in August, where the average value was 2.1 µg/m³ (a factor of 1.2) larger at the Aerosol Station than at the Fonovaya Observatory.

Figure 4. Annually averaged behaviors of mass concentrations of (a) submicron aerosol, (b) black carbon, and their standard deviations at the Aerosol Station (suburban region) and in the Fonovaya Observatory (background) in 2014–2018.

Table 2. Average mass concentrations of submicron aerosol (M_A), black carbon (BC eq), elemental carbon (EC) (µg/m³), and their standard deviation (SD).

| Station                  | Site Type   | Period     | Characteristic | Total Data | Without Fires |
|--------------------------|-------------|------------|----------------|------------|---------------|
|                          |             |            |                | Avg ± SD   | Avg ± SD      |
| Aerosol Station (Tomsk)  | Suburban    | 1997–2018  | M_A, BC eq     | 24.2 ± 15.8| 10.7 ± 2.3    |
|                          |             |            |                | 1.65 ± 0.86| 0.79 ± 0.28   |
| Aerosol Station (Tomsk)  | Suburban    | 2014–2018  | M_A, BC eq     | 24.2 ± 14.9| 20.5 ± 11.9   |
|                          |             |            |                | 1.48 ± 0.9 | 0.62 ± 0.06   |
| Fonovaya Observatory     | Background  | 2014–2018  | M_A, BC eq     | 15.9 ± 5.5 | 12.1 ± 5.0    |
|                          |             |            |                | 0.67 ± 0.06| 0.25 ± 0.07   |
| Novosibirsk              | Urban centre| 2000–2016  | EC             | 5.8 ± 1.8  | 5.3 ± 1.6     |
|                          |             |            |                | 5.3 ± 1.6  | 3.8 ± 2.4     |
| Klutchi                  | Suburban    | 1999–2017  | EC             | 2.7 ± 0.94 | 2.3 ± 0.68    |
|                          |             |            |                | 2.3 ± 0.68 | 1.5 ± 0.6     |
| Zavyalovo                | Background  | 2001–2016  | EC             | 1.3 ± 0.73 | 1.0 ± 0.61    |
| Troposphere (aircraft)   | Background  | 2000–2018  | M_A, BC eq     | 11.8 ± 4.61| 10.9 ± 0.58   |
|                          |             |            |                | 0.58 ± 0.29| 0.57 ± 0.29   |
|                          |             |            |                | 6.9 ± 3.58 | 0.24 ± 0.14   |
|                          |             |            |                | 0.24 ± 0.14| 0.10 ± 0.07   |
|                          |             |            |                | 2.43 ± 1.41| 0.10 ± 0.07   |
|                          |             |            |                | 0.10 ± 0.07| 0.07 ± 0.04   |

Analogous tendencies were also observed for the BC eq concentration (Figures 2b and 4b). The average maximal value in annual behavior over 22 years of measurement (2.7 ± 0.8 mg/m³) was revealed in January, and the minimal was found (0.79 ± 0.15 µg/m³) in June. Over the last 5 years, the maximum in seasonal oscillations increased to 3 ± 1 µg/m³ and shifted toward February, and the minimum of 0.62 ± 0.06 µg/m³ shifted toward May. Under the background conditions in this same period, the maximal seasonal variations in the BC eq concentration were detected in these same months, with values of 1.4 ± 0.3 µg/m³ and 0.25 ± 0.07 µg/m³, respectively. Thus, the maximal contribution of black carbon from urban sources over the period of 2014–2018 at the Aerosol Station was recorded in February—the BC eq concentration was 1.6 µg/m³ larger than at the Fonovaya Observatory, with a minimal difference of 0.37 µg/m³ recorded in May.
Figure 5 presents the seasonally averaged concentrations of EC at three sampling sites in Novosibirsk Oblast. Maximal EC concentrations were observed in the winter season in both Novosibirsk (6.9 ± 2.2 μg/m³) and Klutchi village (3.4 ± 1.2 μg/m³). The minimal values in the urban zone (3.8 ± 2.4 μg/m³) were noted in summer. In Klutchi village, the minimum was in autumn (September-October), with EC concentration values of 1.5 ± 0.6 μg/m³. The seasonal behavior of the concentrations of elemental carbon at three stations in Novosibirsk Oblast almost completely repeated the dynamics of the variations in black carbon concentration in the atmosphere of the Tomsk Oblast. The EC concentration in the atmosphere of the background region (Zavyalovo village) at summertime had been 1 ± 0.61 μg/m³, a factor of 1.5 lower than in Klutchi village, and a factor of 3.8 lower than in Novosibirsk.

![Figure 5. Annually averaged behavior of EC concentration and its SD in Novosibirsk (urban center), Klutchi village (suburban) and Zavyalovo village (background; 2000–2016).](image)

4.1.2. Aircraft Measurements

We note that processing of data from the aircraft measurements of the vertical distribution of the concentrations of aerosol and absorbing substance over 2000–2018 showed that seasonal variations in these quantities fit well into the variability range of sensing results published previously [47]. High surface concentrations of aerosol particles for winter months can be explained quite readily (Figure 6). Obviously, the increased aerosol content at altitudes of 0.5–3 km in summer (with the maximum in August, in our dataset) was caused by the effect of smoke from forest fires, as well as by the increase in the height of the mixing layer as the surfaces warmed. In autumn, the total content tended to decrease, seemingly because of a larger number of days with precipitation and fog, which favored cleaning of the atmosphere and eliminated the effect of the underlying surface.

The maximal mass concentration of submicron particles at the 0.5-km level (15.8 μg/m³) was observed in August, and the minimal was in November. The higher the altitude, the smaller the aerosol concentration. At an altitude of 5 km, the maximum of seasonal oscillations shifted towards the cold period of the year. In February, the aerosol concentration was 2.8 μg/m³, gradually decreasing until reaching a minimum of 0.57 μg/m³ in November.

Regarding the seasonal variations in the black carbon concentration, it should be noted that increased values at the 0.5-km level were manifested in the warm period of the year (with a maximum of 1.15 μg/m³ in September and a minimum of 0.35 μg/m³ in December). The tendency of an increased BC_{eq} concentration was retained up to an altitude of 3 km in the warm period compared with the cold period of the year. At an altitude of 5 km, the seasonal behavior changed, and the BC_{eq} concentration minimum was observed in summer. The black carbon concentration decreased to 0.03 μg/m³. In the cold period, the BC_{eq} concentration grewed at this altitude (the highest value of 0.1 μg/m³ observed in February). It should be noted that separate bursts of submicron particle concentrations were often recorded during the sensing period at different altitudes, with a high content of absorbing substance (Figure 6). Evidently, these particles were anthropogenic in origin [48–52].
To estimate the possible contribution of anthropogenic sources to the state of aerosol in the troposphere, we used the parameter $P = \frac{BC_{eq}}{MA}$, characterizing the “blackening” degree of the submicron particles. In the classification suggested for “aerosol weather” types, a value $P > 0.05$ (or 5%) was characteristic for smog situations, caused by anthropogenic pollutants [28].

Our analysis showed that the average $BC_{eq}$ fraction content in aerosol $P$ was in the range of 0.042–0.098 at all altitudes over the entire period of measurements.

Considering that some ambiguity existed in identifying the smog situations at the boundary separating different “aerosol weather” types according to the criterion ($P = 5\%$), for reliability, we selected situations for which the $P$ value exceeded 7%. It was found that increased (>0.07) $P$ values were observed most often at altitudes of 500 m and 3000–5500 m, and that their frequency of occurrence had a pronounced seasonal behavior (Figure 7). At altitudes from 500 to 5500 m, this was observed in 5–5.5% of situations in the cold period; while in summer months, the number of these situations was 1% of the total data.

Hence, it can be concluded that, on the whole, the contribution of anthropogenic sources is small, and, when creating prognostic models, the average values of the studied characteristics can be quite successfully used to describe the aerosol characteristics for background conditions of the troposphere in the region of Southwestern Siberia.
4.2. Interannual Variations

In the availability of multiyear observations in the troposphere of Southwestern Siberia, it is important to estimate possible long-term tendencies of variations in the submicron particle and black carbon contents. Taking into account the relatively short lifetime of aerosol particles in the lower troposphere, among the main processes causing their interannual variations, we can single out the following factors: changes in the macroscale and mesoscale atmospheric circulations, determining alternation among air masses of different origins and trajectories of their motion, and a growth or decrease in the contribution of anthropogenic sources, located in immediate vicinity of the observation sites. As was shown for the territory of Siberia, it is also necessary to take into consideration the specific contribution of smoke from wildfires. The frequency of their occurrence is known to be determined heavily by both the changes in weather conditions on the whole on vast territories, and the intensifying role of the human factor.

4.2.1. Surface Measurements
Multiyear Behavior of Submicron Aerosol Concentration

As expected, the largest annually average values of submicron aerosol mass concentration at the Aerosol Station were recorded in 1997, 2012, and 2016, i.e., the years with the most severe fires in duration, covering the greatest territory in the Siberian region (Figure 8a). Accounting for the contributions of situations with smoke from forest fires led to an increase of the annual average aerosol concentration in 2012 by 20 μg/m³. A minimal annual average aerosol concentration was observed in 2010. At the same time, our estimates showed that no reliable trend of interannual M₄ variations was present (Table 3). The trend values were calculated from linear regression, constructed using the least squares method. It is noteworthy that there was a significant (at p-value = 0.05) decrease of aerosol concentration by 2.3 ± 1.1% per year only in spring season, and a clarification of the causes for this requires a separate study.

It can be clearly seen that increases in the annual average particle concentrations could be discerned in certain periods, likely associated with changes in the circulation processes under the influence of quasibiennial oscillations (QBO) [53,54].
Figure 8. Multiyear behavior, SD, and trends of annual average mass concentrations of (a) submicron aerosol and (b) black carbon at the Aerosol Station (suburban) in 1997–2018. Blue shadings show SD of total dataset of (a) mass concentrations of submicron aerosol. Gray shadings show SD of total dataset of (b) mass concentrations of black carbon. Open diamonds indicate concentrations without accounting for the contributions from situations with wildfires.

Table 3. Trends (% per year) of the concentrations $M_A$, $BC_{eq}$, EC ($\mu g/m^3$) and their errors and significance ($p$-value), given in parentheses, at different stations.

| Station                  | Site Type | Characteristic | Period       | Year       | Winter  | Spring   | Summer    | Autumn    |
|--------------------------|-----------|----------------|--------------|------------|---------|----------|-----------|-----------|
| Aerosol Station (Tomsk)  | Suburban  | $M_A$          | 1997–2018    | 0.01 ± 0.9 (0.9) | 0.7 ± 0.4 (0.1) | 0.8 ± 0.5 (0.1) | 0.8 ± 0.5 (0.1) | 0.8 ± 0.5 (0.1) |
| Aerosol Station (Tomsk)  | Suburban  | $BC_{eq}$      | 1997–2018    | −1.5 ± 0.6 (0.03) | −1.1 ± 0.5 (0.09) | −3.7 ± 0.6 (0.001) | −0.6 ± 0.1 (0.7) | −3.3 ± 0.8 (0.001) |
| Novosibirsk              | Urban Centre | EC         | 1999–2016    | −3.6 ± 1.3 (0.01) | −0.2 ± 1.5 (0.8) | −6.5 ± 2.5 (0.02) | −5.6 ± 2.6 (0.04) | −1.8 ± 2.3 (0.4) |
| Klutchi                  | Suburban  | EC             | 1999–2017    | −3.0 ± 1.3 (0.03) | −2.1 ± 1.5 (0.2) | −3.4 ± 2.1 (0.1) | −2.4 ± 1.6 (0.1) | −5.1 ± 1.8 (0.01) |
| Zavyalovo                | Background | EC             | 2001–2016    | -          | -       | -        | −5.8 ± 2.5 (0.03) | -        |

Multiyear Behavior of Black Carbon Concentration

An analysis of the entire multiyear time series revealed a reliable decrease in the concentration of absorbing substance in the composition of submicron fraction of particles by $1.5 \pm 0.6\%$ per year, with the strongest reductions in spring and fall, by $3.7 \pm 0.6$ and $3.3 \pm 0.8\%$ per year, respectively (Figure 8b and Table 3). It should be specially noted that the annual average $BC_{eq}$ concentration values were practically at the same level within this time series from 2000 to 2004, followed by a decrease until 2010, then changed to the growth, which was most intense in the winter season [55]. As expected, the maximal annually average $BC_{eq}$ concentrations were characteristic for 2012 data, owing to the anomalously high contribution of wildfires in the summer season. We note that, as in multiyear variations of the total mass concentration of submicron aerosol, in this case, we again discerned quasibiennial oscillations of annual average values.

Multiyear Behavior of Elemental Carbon

Considering the results of observations (1999–2017) of the concentrations of elemental carbon EC in the Novosibirsk oblast, we highlight the main points. The trend of annual average and seasonal mean values was negative over this period at all observation sites (Table 3). Aerosol samples were only collected in summer in Zavyalovo village. There
was a decrease in annual average values of EC concentration in Novosibirsk and Klyuchi village, and in summer values in Zavyalovo village until 2007–2008, observed with a reliable trend of \( -11 \pm 2.3\% \) (\( p\)-value = 0.002) per year in Novosibirsk (Figure 9). Then the decrease changed to an increase in Novosibirsk, and in Klyuchi and Zavyalovo the decrease slowed down.

![Figure 9. Multiyear (1999–2017) behavior of annual average EC concentrations and their trends (solid lines) in Novosibirsk (urban center), Klutchi village (suburban), and Zavyalovo village (background). Dashed lines show EC concentration trends in Novosibirsk during 1999–2007 and 2008–2016.](image)

Multiyear Variations in the Contribution of Anthropogenic Sources

To determine long-term variations in the effect of urban sources on the measured aerosol and black carbon concentrations, we will consider the data, obtained in winter and summer periods (the total dataset was processed to remove situations with the effect of smokes from wildfires).

The aerosol and \( \text{BC}_{\text{eq}} \) concentrations increased in the winter season at the Aerosol Station (suburban); at the same time, no interannual trend existed in this period of the year at the Fonovaya Observatory (background; Figure 10). The difference between the average wintertime values of the aerosol in the suburb and under the background conditions increased from 15 \( \mu \text{g}/\text{m}^3 \) in 2014 to 20 \( \mu \text{g}/\text{m}^3 \) in 2018, i.e., by a factor of 1.3. The divergence in black carbon values between the suburban conditions and the background region during winter increased from 0.94 \( \mu \text{g}/\text{m}^3 \) in 2014 to 1.75 \( \mu \text{g}/\text{m}^3 \) in 2018, i.e., by a factor of 1.9. Thus, from a comparison of the measurements, it can be seen that in the winter season, there was a permanent intensification of the anthropogenic effect of the city on the aerosol composition at a suburban station. In the summer period, the aerosol concentration was found to decrease at both stations, while the \( \text{BC}_{\text{eq}} \) concentration decreased much less at both stations. The absolute differences in the monthly average values between the urban and background conditions, averaged over 5 years of measurements, were from 2 \( \mu \text{g}/\text{m}^3 \) in August to 20 \( \mu \text{g}/\text{m}^3 \) in January for aerosol, and from 0.4 \( \mu \text{g}/\text{m}^3 \) in June to 1.6 \( \mu \text{g}/\text{m}^3 \) in January for black carbon. These estimates show that at the Aerosol Station, the \( \text{BC}_{\text{eq}} \) fraction in aerosol increased at a rate of 6 \( \pm \) 2\% per year, corresponding to the absolute increment in the \( \text{BC}_{\text{eq}} \) fraction \( \Delta P \sim 0.004 \) per year. In turn, no \( P \) increase was observed at the Fonovaya Observatory.
4.2.2. Aircraft Measurements

From our analysis of the multiyear (2000–2018) measurements in the troposphere, we concluded that no reliable changes were observed in the concentration of submicron particles at all altitudes (Figure 11a and Table 4). Of note is the considerable increase in aerosol concentration at small altitudes in 2012, most likely associated with the effect of severe fires that raged in Siberia in that year.

In the time behavior, we can clearly discern a minimum value in 2010. In the interannual dynamics of the $\text{BC}_{\text{eq}}$ concentration in the period until 2010 at all altitudes, there was a decrease in concentration, followed by an increase. As a consequence, black carbon increased at all altitudes over the entire period (2000–2018) of observation. At altitude of 1.5 km over the period of 2000–2010, there was a reliable negative trend of $-7.7 \pm 3.3\%$ per year at the $p$-value = 0.04, and then the $\text{BC}_{\text{eq}}$ concentration started to increase. Over the entire period of 2000–2018, black carbon showed a positive trend of $5.3 \pm 2.2\%$ per year at an altitude of 1.5 km, found to be significant at $p$-value = 0.05 (Figure 11b and Table 4). After 2010, the $\text{BC}_{\text{eq}}$ concentration growth was especially marked at the altitude of 7 km in the period of 2011–2015.

The most characteristic feature of the seasonal variations in black carbon concentrations was their increase in the summer and fall periods at practically all altitudes, with a maximal growth of $9.7 \pm 2.5\%$ per year ($p$-value = 0.003) observed in summer in the mixing layer at an altitude of 0.5 km.
Figure 11. Multiyear behaviors of annual average concentrations of (a) aerosol and (b) black carbon at different altitudes in clear-sky troposphere (Zavyalovo village) in 2000–2018. Solid lines show trends in 2000–2018, and dashed lines show BC$_{eq}$ trends in 2001–2010 and 2011–2015.

Table 4. Trend (% per year) concentrations of MA, BC$_{eq}$, and their errors and significance ($p$-value), given in parentheses, at different altitudes in 2000–2018.

| Characteristic | 500 m | 1500 m | 3000 m | 7000 m |
|----------------|-------|--------|--------|--------|
| MA             | 0.5 ± 1.7 (0.8) | −1.2 ± 2.2 (0.6) | −0.1 ± 2.5 (1) | −1.6 ± 2 (0.4) |
| BC$_{eq}$      | 3.3 ± 2.0 (0.11) | 5.3 ± 2.2 (0.03) | 1.9 ± 2.9 (0.5) | 2.9 ± 2.5 (0.3) |

The data presented here indicate that the lower troposphere of Southwestern Siberia is already beginning to exhibit (still weak) signs of increasing anthropogenic load, probably associated with the developing industry and urban infrastructure.
5. Conclusions

In this work, we analyzed the results from multiyear studies of submicron aerosol mass concentration and black carbon content in the troposphere of Southwestern Siberia. To correctly identify the aerosol states characteristic of the study region, we considered a long observation time series obtained in the surface atmospheric layer at five ground-based stations, as well as the results from regular flights of the aircraft laboratory over background region in Novosibirsk oblast.

The annual average values of the studied parameters were considered in order to estimate the interannual variations. The submicron aerosol and black carbon concentrations in the surface atmospheric layer were found to be maximal in 1997, 2012, and 2016, when the largest number of wildfires occurred in the Siberian region. We revealed that the effect of quasibiennial oscillation of atmospheric circulation manifested in certain periods of their interannual variations. Estimates showed that no reliable trend of interannual variations in $M_A$ existed, but a reliable decrease of $1.5 \pm 0.6\%$ per year was distinctly manifested in the concentration of the absorbing substance in the composition of the submicron fraction of particles, with the strongest decreases in spring ($3.7 \pm 0.6\%$ per year) and fall ($3.3 \pm 0.8\%$ per year).

Synchronous measurements at the Aerosol Station (suburban) and the Fonovaya Observatory (background) over the period of 2014–2018 were compared to estimate the aerosol contribution of the industrial center to the aerosol composition in the surface atmospheric layer. The contribution of anthropogenic sources in the suburban region was shown to be the largest in winter—in January and February, the aerosol concentration in the suburb were on the average $20 \mu g/m^3$ larger than under the background conditions. The minimal difference was observed in the warm period of the year: in August, at the Aerosol Station, the average value was $2 \mu g/m^3$ larger than at the Fonovaya Observatory. The maximal contribution of the city to the black carbon content was recorded at Aerosol Station in February, when the $BC_{eq}$ concentration was a factor of $1.6 \mu g/m^3$ larger than at the Fonovaya Observatory, and the minimal ($0.4 \mu g/m^3$) was found in May. From a comparison of the measurements, it can be seen that in the winter season, there was a permanent intensification of the anthropogenic effect of the city on the aerosol composition at the suburban station. It should be noted that the seasonal behavior of EC at three stations in Novosibirsk Oblast almost completely matched the dynamics of variations in the $BC_{eq}$ concentration in the atmosphere of Tomsk Oblast. The EC concentration in the atmosphere of the background region (Zavyalovo village) in summertime was a factor of 1.5 lower than in suburban (Klutchi village) and a factor of 3.8 lower than in Novosibirsk.

Analysis of the multiyear (2000–2018) measurements in the troposphere revealed no reliable changes in the concentration of submicron particles at all altitudes. In contrast to the mass concentration of the submicron aerosol, the black carbon content was found to show an increasing tendency. In the multiyear behavior of the $BC_{eq}$ concentration at all altitudes, there was a decrease in concentration in the period until 2010, followed by an increase in black carbon. As a consequence, over the entire period (2000–2018) of observations, $BC_{eq}$ increased at all altitudes. The positive trend of $5.3 \pm 2.2\%$ per year at an altitude of 1.5 km was found to be significant at $p$-value = 0.05.

Summarizing the results of our analysis of multiyear studies as a whole, we note that the obtained quantitative values of the aerosol characteristics, and the resulting seasonal behavior and trends of their interannual variations, are necessary for their use as input parameters for regional-scale prognostic models.

Author Contributions: Conceptualization, M.P.; instrumentation maintenance, V.S.; measurements, D.C., S.P., and E.Y.; calculations: E.Y., D.C., and S.P.; analysis of results, M.P., V.K., VM., E.Y., D.C., and S.P.; writing original draft, E.Y. and S.P.; writing review and editing, M.P. All authors have read and agreed to the published version of the manuscript.
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Funding: The long-term measurements were supported by the Ministry of Science and Higher Education of the Russian Federation (IAO SB RAS). The analysis was carried out in 2020, with the financial support of the Russian Science Foundation (agreement No. 19-77-20092).

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: Data availability. Data can be obtained from Mikhail Panchenko (pmv@iao.ru) upon request.

Conflicts of Interest: The authors declare no conflict of interest.

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