A Decade of Aerosol Optical Properties Measurements over Athens, Greece

Ioannis-Panagiotis Raptis 1,*, Stelios Kazadzis 2, Vassilis Amiridis 3, Antonis Gkikas 3, Evangelos Gerasopoulos 1 and Nikolaos Mihalopoulos 1,

1 Institute for Environmental Research and Sustainable Development, National Observatory of Athens, GR15236 Athens, Greece; egera@noa.gr (E.G.); nmihalo@noa.gr (N.M.)
2 Physicalisch-Meteorologisches Observatorium Davos, World Radiation Center, CH-7260 Davos, Switzerland; Stelios.Kazadzis@pmodwrc.ch
3 Institute for Astronomy, Astrophysics, Space Applications and Remote Sensing, National Observatory of Athens, GR15236 Athens, Greece; vamoir@noa.gr (V.A.); agkikas@noa.gr (A.G.)

* Correspondence: piraptis@noa.gr

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Abstract: Long-term ground-based measurements of aerosol optical properties in Athens, Greece, for the period 2008–2018 performed by the National Observatory of Athens are used in order to investigate the aerosol climatology of the area. In this study, we utilize quality-assured measurements of the aerosol optical depth (AOD), Single Scattering Albedo (SSA) and Ångström exponent obtained by CIMEL photometers in the framework of the Aerosol Robotic Network (AERONET) to extract the seasonality and the trends of aerosols in the region. Higher aerosol loads are found during spring and summer months. A 1.1% per year decrease for AOD at 440 nm and 0.4% decrease per year for SSA during the studied period are recorded. Collocated and synchronous PM$_{10}$ values, for a five-year period, are used in order to study ground-level conditions. Also, the Planetary Boundary Layer Height from ERA-5 is used to investigate the stratification of the particles. The classification of aerosols using AERONET data is performed to separate dust, biomass burning, polluted urban, marine and continental dominant aerosol mixtures. Also, the characterization of AOD provided by Copernicus Atmosphere Monitoring Service (CAMS) is investigated. Finally, seasonal AOD trends recorded from AERONET from satellite sensors (MODIS-Aqua/MODIS-Terra) and estimated by CAMS are examined, and significant differences have been found.

Keywords: aerosols; climatology; AERONET; Athens; MODIS; aerosol classification; aerosol trends

1. Introduction

Aerosols play an important role in Earth’s climate system, as has been reported by the Intergovermental Panel on Climate Change (IPCC) [1], and also affect human health in urban and agricultural areas [2–4]. Aerosols absorb and scatter solar radiation; however, these mechanisms still remain a major source of uncertainty in climate modelling [5]. Additionally, aerosol estimation is important for diverse applications such as in the correction of the surface’s satellite retrievals [6–8] and in forecasting solar energy [9,10].

Over the last two decades, significant improvements have been performed for surface networks and the satellite retrievals of aerosols, contributing to major improvements in the monitoring and understanding of the related procedures. The major surface aerosol networks include AERONET, which operates hundreds of stations around the globe [11]; SKYNET, which operates more than 60 stations globally with main focus on Asia [12]; and the Global Atmospheric Watch/Precision Filter Radiometer (GAW/PFR), which operates approximately 20 stations with long-term measurements [13,14]. Various homogenization
activities for these different networks have been reported [15,16]. Data from these networks have been used frequently for climatological studies. Satellite aerosol retrievals are available from several instruments [17], such as Microwave Integrated Retrieval System (MISR) [18], Cloud-Aerosol Lidar and Infrared Pathfinder Satellite Observations (CALIPSO) [19], Advanced Along-Track Scanning Radiometer (AATSR) [20]. Satellite-based products of aerosol properties, although geographically denser, are however more uncertain than ground-based measurements.

Aerosol trends in urban environments have attracted specific interest over recent years as anthropogenic emissions (traffic, industry) undergo significant interannual changes. In addition, satellite retrievals above densely populated areas are subject to significant errors [17], leading to different indications of trends in a number of areas when satellite-derived time series are used [21]. Model sulfate concentrations show a slower rate of decline than sulfur emissions across Western Europe and the USA, mainly as a result of the increased availability of H$_2$O$_2$ in clouds, which alters the oxidation rate of SO$_2$ [22]. In Europe, an average aerosol decrease of 2.3% year for the period 1995–2017 has been reported by the Aerosol, Clouds and Trace Gases Research Infrastructure (ACTRIS) [23].

The current study focuses on the city of Athens, Greece, which is located near the coastline in the eastern Mediterranean. The Athens greater area constitutes a complex aerosol environment with a multitude of natural and anthropogenic sources. A common natural procedure in the area is the emission of marine aerosols by the sea surface, which are advected by the wind speed/sea waves and then chemically processed by interactions with gases and water vapor [24]. The long-distance transport of Saharan desert dust aerosols is occasionally observed, especially in spring/summer months, with high aerosol loads in the free troposphere in most cases [25–27]. Marinou et al. [28] and Koukouli et al. [29] found a very small negative trend in the dust optical depth in the eastern Mediterranean for the period 2007–2015 using satellite data. Wildfires also contribute to the aerosol mixture in the area occasionally, either by nearby events [30,31] or by long-range transport [32,33]. Athanasopoulou et al. [31] showed that wildfires in Greek forests could lead to up to three times higher aerosol optical depth (AOD) values in the area during summer. The Mediterranean basin is moderately to highly polluted over summer and significantly lower during winter months because of air masses originating from Turkish cities and European industrial areas [34]. A major local aerosol source in the greater area of the Athens basin is biomass burning [35–39], which has significantly increased in winter months since 2010 due to the economic crisis and the associated wood burning for residential heating. In addition, secondary aerosol formation makes a significant contribution in the area [40–44]. Especially during summer months, when the long sunshine duration enhances photochemical activity, secondary aerosols are dominant in the area, particularly when air masses arrive from the northern parts of the Balkan peninsula [45]. Emissions of aerosols linked to vehicle traffic have decreased, but the trend has changed since 2014, which is associated with the lifting of the ban of diesel vehicles in the metropolitan area [44]. An estimate of the contribution of the different sources to the average AOD loadings over Athens is provided by Gerasopoulos et al. [25], who reported that the annually averaged AOD is 40% due to regional and local sources, 23% due to dust from the Saharan desert and 22% due to European continental sources.

In the present study, we have exploited a 10-year-long data set recorded by the Aerosol Robotic Network (AERONET) in the city of Athens. The seasonality of the aerosol optical properties is investigated and discussed. A hybrid database that combines columnar variables from AERONET and in-situ PM$_{10}$ measurements has been used as a proxy for the layering of aerosols as well as in respect to the Planetary Boundary Layer Height (PBLH). The classification of aerosols using well-established approaches and modeled data is presented and discussed. Finally, trends in optical parameters, both using AERONET data and modelled data, are compared.
2. Data and Methodology

2.1. Theoretical Background

Aerosol optical depth is the quantity that describes the attenuation of radiation when propagating in an aerosol layer and indirectly quantifies the presence of aerosols in an atmospheric column. It is defined as follows:

$$\text{AOD}_\lambda = \int_0^{\text{TOA}} b_{\text{aer}}(\lambda, h) \, dh$$

(1)

where $\lambda$ is the wavelength, TOA is the top of the atmosphere and $b_{\text{aer}}$ is the extinction over a vertical optical path. Additionally, Ångström [46] proposed the following formula, which includes the spectral dependence of the extinction:

$$\text{AOD}_\lambda = \beta \lambda^{-\alpha}$$

(2)

where $\alpha$ is the Ångström exponent, which parameterizes the spectral dependence of the extinction, and $\beta$ is a variable indicative of the number of particles found in the solar beam path. Using solar direct irradiance measurements from the ground, AOD is calculated through the Beer–Lambert law, which, when solved for atmospheric boundary conditions, is expressed as follows:

$$I_\lambda = I_{0,\lambda} \exp(-m\tau) R^{-2}$$

(3)

where $I_\lambda$ is the irradiance measured at ground level at wavelength $\lambda$, $I_{0,\lambda}$ is the extraterrestrial irradiance at wavelength $\lambda$, $m$ is the optical air mass along the line of sight connecting the observations point and the sun, $\tau$ is the total optical depth and the term $R$ is the Sun–Earth distance in astronomical units, normalized to the variations around the mean distance. Then, the AOD is extracted from $\tau$ by eliminating any other extinctors including trace gases and physical processes such as Rayleigh scattering.

The Ångström exponent $\alpha$ is usually calculated as the ratio of AOD between two wavelengths $\lambda_1$ and $\lambda_2$:

$$\frac{\Delta \text{AOD}_{\lambda_1}}{\Delta \text{AOD}_{\lambda_2}} = \left(\frac{\lambda_1}{\lambda_2}\right)^{-\alpha}$$

(4)

The Single Scattering Albedo (SSA) is a variable that is defined as the ratio of scattering to total extinction. The SSA at a wavelength $\lambda$ is defined as the relative contribution of the aerosol scattering extinction ($b_{\text{sca}}$) to total extinction ($b_{\text{sca}} + b_{\text{abs}}$), thus indirectly also describing the absorption contribution ($b_{\text{abs}}$):

$$\text{SSA}(\lambda) = (b_{\text{sca}}(\lambda))/(b_{\text{sca}}(\lambda)+b_{\text{abs}}(\lambda))$$

(5)

The values of SSA could theoretically range from 0 to 1, but for aerosol layers in the Earth’s atmosphere, they are very close to 1, as aerosols mainly scatter the solar light, and are rarely found to be lower than 0.65 [47], when very absorbing aerosols dominate the layer.

The planetary boundary layer is the lowest part of the troposphere, is directly affected by the surface conditions and responds to surface forcing in very short time scale [48]. The planetary boundary layer height (PBLH) is the height that distinguishes the atmospheric boundary layer from the free troposphere, and the highest values on cloudless days are detected around local noon. On cloudy days, the PBLH has a more complex behavior, resulting in a complicated stratification. In most cases, local emissions of aerosols and trace gases are trapped in this layer; thus, it severely affects the dispersion of pollutants. Stratification in the lower troposphere is the main factor leading to the fact that the columnar properties of aerosols are often different from those measured at the surface. Hence, PBLH could be used to interpret patterns caused by the stratification.

When addressing air quality and public health issues, instead of the columnar AOD, the concentration of different fractions of particulate matter (PM) at ground level is used, with $\text{PM}_{10}$ being the most traditional and most commonly used fraction in terms of legislation obligations.
PM$_{10}$ is defined as the concentration of all solid or liquid-suspended particles that have diameters less than 10 µm. PM$_{10}$ can be monitored by using different methods such as sampling techniques on filters (reference method)—e.g., cascade impactors which use size selective inlets that favor either a certain particle size range or a particular upper cut-off point—or (almost) real-time techniques such as tapered element oscillating microbalance (TEOM) and light scattering systems—e.g., optical particle counters—while currently the most common method is beta gauge monitors (using the attenuation of beta radiation through a filter of specific particle loading).

2.2. Site

Athens is a city of 3.7 million inhabitants with an estimated 2.5 million automobiles and heavy traffic, which represents the single greatest source of local emissions. The area has been heavily deindustrialized over recent decades, but there are still emissions linked to factories and fossil fuel. Athens is located in a basin in which three mountains with a height of around 1 km trap most of the urban emissions in the greater area, causing poor ventilation. The dispersion of the pollutants is mainly caused by the sea breeze along the NE–SW axis [49]. Additionally, there are common cases of the long-distance transport of air masses from arid areas of Northern Africa, frequently associated with dust events that affect the area [50,51]. Athens has a temperate climate with warm and dry summers and wet and mild winters, which is typical for the Eastern Mediterranean. The measurements used in this study were conducted at the urban background site of the National Observatory of Athens (NOA) at Thissio (37.97° N, 23.72° E), which is located at an elevation of 130 m a.s.l. and 8 km from the coastline, in a moderately populated area, where the influence of direct local emissions is limited. A detailed description of the measuring site can be found in Paraskevopoulou [35].

2.3. Measurements

Data from the AERONET station ATHENS-NOA have been used for the period May 2008–September 2018. The instrument is located at the site described in the previous paragraph, and the horizon view is clear at a 360° viewing angle.

The CIMEL Sun-photometer is a filter radiometer developed by Cimel Electronics (Paris, France), which performs direct Sun and sky radiance measurements. Measurements are performed at nine bandpass filters between 340 and 1640 nm (eight of them are dedicated to AOD retrieval, and one us used for integrated water vapor (IWV)). During this time period at the station CIMEL CE318, photometer #440 operated as the main instrument, with some months of operations of CIMEL CE318 photometers #110, #240 and #395 as replacement instruments. Instruments #110 and #240 recorded only four channels (440, 675, 870 and 1020 nm), which shortened the time series of the other wavelengths by a total of 350 days (combined for these instruments). Additionally, gaps in the time series are expected due to the frequent calibrations (annually) demanded by the network protocol. All calibrations were performed at Laboratoire d’Optique Appliquée (LOA) at Université de Lille, France.

Direct measurements are performed usually every 10–15 min. These measurements are processed centrally and are widely available from the Aerosol Robotic Network (AERONET) [11]. Aerosol optical depth data are computed for three data quality levels: Level 1.0 (unscreened), Level 1.5 (cloud screened), and Level 2.0 (cloud-screened and quality-assured). Thus, gaps in time series are caused by cloudy skies and multi-day gaps by instrument malfunction. Additionally, there is an around 2-month-long gap in the time series for every calibration of the instrument.

In this study, AERONET version 3 retrievals have been used [52,53] at level 2.0 for direct Sun and inversion products to guarantee the highest quality. The unique exception is Single Scattering Albedo (SSA) inversion retrievals, for which level 1.5 was used. This practice is relatively common in climatology studies, because AERONET level 2.0 criteria for this product are very strict (Solar Zenith Angle (SZA) > 50° AOD > 0.4) and for many areas, including Athens, such cases are rare. For the presented dataset, level 2.0 filtered out all but 186 SSA level points in 10 years, while at level 1.5, there are 8933 measurements available. A conditional sampling has been applied to level 1.5 data
in order to ensure that the quality assurance filtered out only based on AOD values. The sampling criterion applied in the current dataset has been described by Kazadzis et al. [54]. The usage of level 1.5 should be performed while always taking into account the higher uncertainty of this product, which could be up to 0.04 higher than level 2.0 depending on the AOD and SZA.

The monitoring of in situ PM$_{10}$ takes place continuously at Thissio from NOA as part of the routine station measurements, and data are available from 1 January 2014. The PM$_{10}$ concentrations near the surface were measured by means of a beta-attenuation monitor (Eberline, FH 62 I-R), which pulls ambient air through the sampling line, and a size-selective inlet for PM$_{10}$, which deposits particles on a clean section of a filter tape (single filter spot principle) and utilizes the radiometric attenuation (Kr-85 noble gas, with a maximum energy 0.67 MeV) by a two-beam compensation method [55]. Measurements are stored at an initial, 1 min temporal resolution; in this study, we have used the hourly averages. The instrument is located 50 m away from the AERONET site.

The Copernicus Program and its Atmosphere Monitoring Service (CAMS) combines state-of-the-art atmospheric modelling of aerosols with Earth observation data to provide information services covering European air quality, global atmospheric composition, climate, and UV and solar energy [56]. CAMS is part of European Centre for Medium-Range Weather Forecasts (ECMWF) numerical weather prediction and combines the assimilation of satellite data on aerosols and trace gases with chemical modelling and ground-based measurements [57,58]. CAMS is one of the main tools used in Europe for forecasting air quality at regional scales. Emissions sources are also included in the modelling procedure, as well as sedimentation and wet and dry deposition processes [59]. In the current study, we have used 3 h time step outputs from CAMS for pixels corresponding to instrument locations, for the variables AOD at 550 and 1020 nm and for dust, black carbon, organic matter, sea salt and sulfate AOD for the period 2008–2018. Sulfate is assumed to be non-absorbing, and all the assumptions used could be found in the work by Benedetti [59]. The spatial resolution of CAMS data is about 80 km, which could be a source of inconsistencies when compared with a point Sun-photometric measurement, especially in a complex environment such as Athens. The CAMS dataset has potential use for climatological studies in areas without ground-based measurements. Thus, we performed an analysis of this data, despite the expected differences from AERONET, in order to assess the provided climatology and especially the classification of optical depths for different aerosol types.

Spaceborne observations of aerosol optical depth at 550 nm (AOD550), acquired from the Moderate Resolution Imaging Spectroradiometer (MODIS), have been utilized in order to investigate their consistency against ground-based data obtained by the AERONET ATHENS-NOA station. MODIS, mounted on NASA’s twin polar-orbit satellites Terra and Aqua, provides pre-noon (Terra) and post-noon (Aqua) columnar aerosol observations, almost on a daily basis, thanks to its wide swath (~2330 km). The MODIS AOD product analyzed here resulted from the merging [60] of the corresponding retrievals obtained from three independent algorithms applied over dark continental [61,62] and maritime [62,63] surfaces (Dark Target) as well as over land areas via the Deep Blue approach [64]. In the current study, the Level 2 (L2) MODIS AOD values (10 km x 10 km spatial resolution at nadir view), provided by the latest version of the applied retrieval algorithms (Collection 6.1, C061), are used. The aforementioned data are organized in bands (segments of 5 min intervals) and have been downloaded from the Level 1 and Atmosphere Archive and Distribution System (LAADS) Distributed Active Archive Center (DAAC) [65]. In order to ensure the best quality of AOD retrievals, only those associated with a Quality Assurance flag equal to 3 have been processed [66]. MODIS AODs satisfying the quality control criterion are spatially averaged in a circle centered on the Athens AERONET station with a radius of 20 km. All the regional AOD averages are calculated for every day with available satellite data during the period 2008–2018.
3. Results

3.1. Seasonality

Since the aim of this work is to define the climatology of aerosols in Athens area, the monthly means of AOD at all seven recorded wavelengths are presented in Figure 1. Monthly mean values are counter-proportional to wavelength, as 870, 1020 and 1640 nm show very low mean values (almost always < 0.1), while UV wavelengths record the highest values. Retrievals at 1640, 1020, 870 and 675 nm have peak monthly AOD values in April–May, while 440 nm and UV AODs demonstrate the highest values in July–August. AOD values at 675, 870, 1020 and 1640 nm show a secondary peak during summer months. At all wavelengths, the lowest monthly values are recorded in January and December. Additionally, it should be noted that there is a local maximum at all wavelengths for November mean values. This behavior is mainly explained by the maximum values recorded during November 2010 and 2011. It should be noted that during November 2011, there was a severe drought in Eastern Europe, which resulted in wildfires in Ukraine, Moldova and Slovakia. It has been reported in earlier studies that, depending on synoptic conditions, plumes from these areas reach Greece [67].

![Figure 1](image_url)  
**Figure 1.** Monthly mean of aerosol optical depth (AOD) at all seven Aerosol Robotic Network (AERONET) wavelengths, calculated for months with at least 15 days of data.

Monthly values of the Ångström exponent of 440–870 nm (Figure 2) show that the AOD peak in July–August is related to higher α values, which imply more fine aerosols in the mixture. Meanwhile, in April–May, the Ångström exponent has the lowest values, which indicates a dominance of coarse mode particles. This attribute explains the different spectral behavior of monthly mean values in Figure 1, where higher values are found at shorter wavelengths with different seasonality. Additionally, it should be highlighted that for January and December, when AOD is generally very low, the Ångström exponent has different behavior. The mean December Ångström exponent is high (1.52), but for January, the mean value is 1.02, which implies almost equal fine and coarse fractions. Figure 2 also reports the number of data points per month in the whole measuring period. Since measurements depend on direct solar irradiance, the amount of data is related to the sunshine duration. January and December have the least data, with 2127 and 2100 points, respectively, and August is the month with the largest dataset, with 13,182 data points.
Figure 2. Monthly mean of AOD 440 nm and Ångström exponent 440–870 nm. The lower plot shows the amount of data per month.

Figure 3 shows the mean size distribution per month, providing more evidence for the arguments presented in the previous paragraph. The coarse mode has its maxima in May, June and April, which is the period with the highest occurrence of Saharan dust events in the area due to enhanced cyclonic activity [50]. Meanwhile, the fine mode shows maximum values during August, September and July. December is the period in which fine mode dominates, while the coarse mode has very low values. In January, the two fractions are almost equal. February also shows a coarse mode double the fine mode. Regarding the coarse mode, June, August and September have their peaks in a higher radius than the other months, suggesting the presence of even bigger particles in the mixture. It should be highlighted that direct sun and inversion retrievals are extracted from different measurements in the data set (direct sun and almucantar), but in the current dataset, the fine/coarse behavior converges.

Figure 3. Mean size distribution per month with AERONET level 2.0.
Mean monthly SSA values at four wavelengths are presented at Figure 4. The lowest values are retrieved during January, July and December; thus, more absorbing particles are presented in these periods. SSA values at 440 nm show very little seasonal variation, ranging between 0.91 and 0.93. This spectral seasonal pattern results in four months (April, May, June and November) with a lower SSA of 440 nm. For the rest of the year, SSA at 440 nm is higher, showing a decrease to longer wavelengths. Thus, the spectral behavior of SSA indicates that, in these periods, the particles also have a varying spectral absorption. Brown carbon is known to highly absorb in UV region, while it is almost non-absorbing above 700 nm [68], which was found in an earlier study to have the highest concentrations in Athens during the same four months [54]. Also, dust aerosols—especially those containing hematite—are known to absorb more at lower wavelengths [69], which also contributes to this behavior for the April–June period.

Figure 5 shows all SSA values at 440 nm against an Ångström exponent of 440–870 nm and respective AOD loads according to colors. SSA retrieval has the highest uncertainty when AOD < 0.2; thus, these data points should be treated with caution, especially when very low SSA values are retrieved. Very high AOD (>0.7) cases could by separated into two different groups: one with high SSA, where scattering particles dominate, and one with lower SSA values and an Ångström exponent < 0.6—a condition which best describes the Saharan dust events in the area. The class of 0.4 < AOD < 0.7 AOD shows a high concentration around the mean SSA values for the period which is equally spread in high and low Ångström exponents. Only 9% of cases in this class have an SSA lower than 0.9. The class of 0.2–0.4 AOD also has a considerable SSA uncertainty, containing a large number of measurements which were rejected at AERONET level 2.0. Thus, there is a number of cases with an uncommonly low SSA (13% lower than 0.85), which should be attributed to the uncertainty.
The comparison between these datasets is considered synchronous; otherwise, daily average values are presented in order to clarify the seasonal variations of the conditions.

3.2. Connection of AOD with PM$_{10}$

For the comparison of in-situ data with AERONET measurements, a new dataset has been created by the linear interpolation of PM$_{10}$ hourly averaged values to the CIMEL recording moments. The comparison between these datasets is considered synchronous; otherwise, daily average values have been compared. In Figures 6 and 7, blues lines represent the mean AOD at 440 nm and mean PM$_{10}$, respectively, for that period. At Table 1, the relative frequencies of the four quadrants created by the mean values (blues lines) are presented in order to clarify the seasonal variations of the conditions.

![Figure 5](image-url)  
**Figure 5.** SSA 440 nm in respect to the Ångström exponent 440–870nm, categorized according to the AOD for the whole period.

![Figure 6](image-url)  
**Figure 6.** AOD at 440 nm plotted against synchronous particulate matter (PM)$_{10}$, separated by season (color), with blue lines for the mean AOD and mean PM$_{10}$. 

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Table 1. Occurrence of cases in each quadrant per season (December-February, March-May, June-August, September-November) for all synchronous values. Quadrants are separated by the mean AOD 440 nm and PM$_{10}$ values for the whole dataset.

<table>
<thead>
<tr>
<th>Quadrant</th>
<th>DJF</th>
<th>MAM</th>
<th>JJA</th>
<th>SON</th>
<th>All Seasons</th>
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<tbody>
<tr>
<td>Lower Left</td>
<td></td>
<td></td>
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<tr>
<td>Low AOD–Low PM$_{10}$</td>
<td>39.5%</td>
<td>29.4%</td>
<td>31.9%</td>
<td>51.6%</td>
<td>42.5%</td>
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<tr>
<td>Lower Right</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Low AOD–Low PM$_{10}$</td>
<td>8.1%</td>
<td>16.7%</td>
<td>23.8%</td>
<td>14.5%</td>
<td>18.9%</td>
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<td>Upper Right</td>
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<tr>
<td>High AOD–High PM$_{10}$</td>
<td>15.1%</td>
<td>35.0%</td>
<td>37.5%</td>
<td>24.2%</td>
<td>23.6%</td>
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<td>Upper Left</td>
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</tr>
<tr>
<td>Low AOD–High PM$_{10}$</td>
<td>36.6%</td>
<td>18.9%</td>
<td>6.90%</td>
<td>9.7%</td>
<td>14.9%</td>
</tr>
</tbody>
</table>

It appears that in winter months, AOD is usually low, but PM$_{10}$ fluctuates, with both low and high values (compared to the annual mean), which indicates a strong stratification, trapping most of the emissions in lower troposphere. This condition is associated with urban emissions (traffic) and especially with heating. The condition of low AOD and high PM$_{10}$ is significantly less frequent in other seasons. Only in summer months is there a high occurrence of the condition of low PM$_{10}$ and high AOD (23.8% of the cases), which indicates high aerosol loads outside the surface layer. One of the factors contributing to this is the presence of dust layers from the Sahara only at higher levels, which occurs mainly during that season, while in the rest of the year, dust appears to be more distributed in height, drastically affecting the layers near the surface [70]. In general, during winter months, 76.1% of the cases have an AOD lower than the annual mean value. Also, summer is the season in which PM$_{10}$ is lower than the annual mean in 66.1% of the cases. Thus, there is a seasonal anti-correlation of low values of AOD and PM$_{10}$. Extremely high values of PM$_{10}$ (>150 µgr/m$^3$) are only recorded during the March–August period and are synchronous at 90.3% with AOD values higher than the annual mean and at 74.5% with AOD values higher than 0.4. Finally, cases with extremely high AOD (>0.8) are separated into two classes: one with high PM$_{10}$ (>150 µgr/m$^3$), consisting of 48.2% of cases, and one with average PM$_{10}$ (23–62 µgr/m$^3$), with 51.8% of cases. All the extremely high AOD cases occur in the March–August period. Thus, extremely high AOD values are always associated with long-range transport and high aerosol concentration at higher layers, mainly dust from Sahara, and in almost half the cases co-occur with high PM$_{10}$, which could be explained either by strong deposition or the presence of an independent layer near the surface. PM$_{10}$ values during winter show maxima in Athens during night-time, when there are no AERONET measurements, but these data are not used in the above-described comparison.

In Figure 7, daily mean values of AOD at 440 nm and PM$_{10}$ have been plotted with respect to PBLH at 12:00 UTC for all available days. PBLH is extracted from the ERA5 database, generated using the Copernicus Climate Change Service Information [71], which uses assimilated values from ECMWF outputs. Cases with low PM$_{10}$ and low AOD values are more frequently (52.1% of cases) linked with days with low PBL, indicating that the majority of aerosols are found in the lower layer and hardly any particles above that. Cases with low AOD but high PM$_{10}$ concentration are usually (88.9% of cases) associated with low BL heights. In theory, for these cases, it is expected that most aerosols are concentrated in a relatively short surface layer and originate from local pollution sources. Cases of low PM$_{10}$ and high AOD are usually measured when PBLH is more than 600 m (84.2% cases). This condition is expected to occur when most particles are found in layers above the PBL. Specific interest should be attracted to cases of high PM$_{10}$ loads and high AOD values for which the PBL height is lower than 600 m, which gives evidence of the existence of two different aerosol layers at different heights. Taking into account the layering of aerosols in urban areas such as Athens is extremely important for several applications, including in health-related areas.
which means these are cases of high coarse aerosol loads of a scattering nature spread across the
atmosphere. Beyond that fact, the classes of 40–60 µg/m³ have statistically higher fine mode fractions. SSA error bars
87.55% of the data with AOD values
brown carbon during winter. Lower
µ the range of 50–60
aerosol dominance. The lowest mean SSA values are found when PM
concentration classes. A high PM
concentration dominance.

Figure 8 shows the mean SSA 440 nm values and mean Ångström exponents according to PM₁₀ concentration classes. A high PM₁₀ (>50 µg/m³) is linked with columnar mixtures with coarse mode aerosol dominance. The lowest mean SSA values are found when PM₁₀ concentrations are recorded in the range of 50–60 µg/m³, which indicates the presence of dust cases in this class and probably some brown carbon during winter. Lower α values are found mainly with high PM₁₀. It is interesting that 87.55% of the data with AOD values > 0.6 also have PM₁₀ values > 60 µg/m³, α < 1 and SSA > 0.9, which means these are cases of high coarse aerosol loads of a scattering nature spread across the column. PM₁₀ classes lower than 40 µg/m³ are associated with higher Ångström exponents; hence, lower concentrations of surface particles have statistically higher fine mode fractions. SSA error bars represent 1σ, which for all PM₁₀ classes covers the largest part of the expected SSA range in the atmosphere. Beyond that fact, the classes of 40–60 µg/m³ have a systematically more absorbing columnar behavior.

Figure 8. Mean SSA against the Ångström exponent for different PM₁₀ clusters (colors); error bars represent 1σ.
3.3. Particles Classification

In Figure 9, the commonly used AERONET aerosol classification described by Dubovik et al. [72] is visualized. This approach utilizes the inversion products of AERONET and, by using some data from stations where the dominant aerosol type is well known, it defines threshold values for each class. The disadvantage of this approach is the precision of the threshold values in cases of complicated aerosol mixtures. It should be highlighted that columnar atmospheric mixtures in urban areas such as Athens can almost never have a single type of aerosols. Thus, the classification should be treated as the characterization of the prevailing type(s) to which the recorded optical properties are closest.

Using this classification, it appears that polluted-type aerosol mixtures are more commonly detected as the characterization of the prevailing type(s) to which the recorded optical properties are closest. Athens can almost never have a single type of aerosols. Thus, the classification should be treated as Athens can almost never have a single type of aerosols. Thus, the classification should be treated as the characterization of the prevailing type(s) to which the recorded optical properties are closest.

When this characterization is applied to the synchronous PM$_{10}$ data, it appears that marine and continental aerosol cases are usually associated with lower PM$_{10}$ values, while biomass burning and dust particles are more frequently recorded with higher PM$_{10}$ values. In Figure 10, the normalized distribution of the classification of the cases in respect to PM$_{10}$ is presented. It is clear that, above 40 µg/m$^3$, the recorded columnar condition is at 82% related to dust or biomass burning. Since these conditions are usually attributed to the long-range transport of aerosols, it is interesting that it also leads to higher concentrations on the ground, indicating that there is at least a partial deposition effect.

![Figure 9](image_url)

**Figure 9.** AOD at 440 nm against the Ångström exponent, classified according to Dubovik et al. [72] for the whole dataset (left). Pie chart of the Dubovik et al. [72]. classification for the whole dataset (right).
CAMS data are used in order to benefit from the classification made by the model, always keeping in mind the considerable differences from measurements. CAMS characterizes the aerosol types by using the emission inventories in combination with meteorological and chemical modelling. Even with extrapolated data at the AERONET wavelength (440 nm), there are considerable differences \((R^2 = 0.74)\) between the AOD values in the datasets. The seasonality of AOD at 550 nm from the CAMS database appears to have the same pattern as AERONET at 440 nm, as shown in Figure 11. It appears in this dataset that there is a steady sulfate aerosol background, fluctuating at around 0.1 AOD units year-round. Dust aerosols show a peak in the period March–June. It is interesting that the dust AOD seasonal pattern is exactly in phase with AOD at 1240 nm and highly correlated to the AOD 550 nm seasonal pattern. It appears that, in the CAMS dataset, the seasonality of AOD should be mainly attributed to dust events. Organic matter aerosols appear to have a peak during July–August, which is probably linked with wild forest fires. Sea salt AOD has lower values during the summer months and is higher from December to March, which is linked to the higher winds during this period. This characterization of AOD is not directly comparable with the earlier classification which labelled the columnar mixture according to optical properties, while here it is based on emissions and modelling. The separation of AOD made by CAMS converges with AERONET retrievals in the detection of frequent dust presence for spring months and the dominance of fine particles during summer.

**Figure 10.** Normalized distribution of cases in each class in respect to PM\(_{10}\) synchronous recordings according to Dubovik et al. [72].
AOD (through the fine mode fraction) and the size of the fine aerosol using this approach is presented linked with anthropogenic emissions. that these cases are dominated by the coarse mode, which is most likely to be linked with dust events; cluster in two regions of the plot, thus having two different typical behaviors: one is associated with the coarse mode, which is most likely to be linked with dust events; the other one is associated with an average fine mode Rf of 0.20 μm and a contribution of the fine mode at less than 30%, meaning that these cases are dominated by the coarse mode, which is most likely to be linked with dust events; the other one is associated with an average fine mode Rf of 0.11 μm and a contribution of the fine mode at more than 70%, meaning that these cases are dominated by the finest particles and are most likely to be linked with anthropogenic emissions.

Figure 11. Copernicus Atmosphere Monitoring Service (CAMS) mean monthly values for AOD at 550 and 1240 nm and AOD for aerosol types (black carbon—BCAOD, dust—DUAOD, organic matter—OMAOD, sea salt—SSAOD, sulfate—SUCAOD).

The Gobbi plot [73] is an approach to gain information regarding the spectral variation of aerosol optical properties (Figure 12). To achieve this classification, the Ångström exponent difference, defined as δα = α(440, 675) − α(675, 870) is used. A visual estimation of the contribution of fine aerosol to the AOD (through the fine mode fraction) and the size of the fine aerosol using this approach is presented in Figure 12. It should be mentioned that low aerosol loads (AOD < 0.1) propagate high uncertainty to α and δα, at up to 50% [50]; thus, the yellow points in this plot should be considered as highly uncertain. The average fine mode Rf is found at 0.13 μm, and the same average is found for the most usual AOD class (0.1–0.4) and for the cases of more than 70% fine mode contribution. Higher AODs appear to cluster in two regions of the plot, thus having two different typical behaviors: one is associated with an average fine mode Rf of 0.20 μm and a contribution of the fine mode at less than 30%, meaning that these cases are dominated by the coarse mode, which is most likely to be linked with dust events; the other one is associated with an average fine mode Rf of 0.11 μm and a contribution of the fine mode at more than 70%, meaning that these cases are dominated by the finest particles and are most likely to be linked with anthropogenic emissions.

Figure 12. Gobbi plot for daily-averaged AERONET data [72]. Black lines represent the size of the fine mode Rf, and cyan lines represent the fixed fraction contribution of the fine mode to total AOD. Colors of dots represent ranges of AOD at 440 nm.
3.4. Long-Term Trends

Figure 13 reports the trends of AOD during the measured 10-year period (2008–2018). Deseasonalization is calculated by eliminating the mean monthly value of the corresponding month. Afterwards, the trend calculations are performed by estimating a linear fit in the deseasonalized timeseries. AOD at 440 nm decreases during the measurement period, at a rate of 1.1% per year. The Greek Ministry of Environment and Energy, in its annual report, has reported a 1.6% decrease of mean PM$_{10}$ daily values, for the same period, for the station Aristotelous, which is located 1 km from NOA's measuring site [74]. Thus, the negative trend is mainly due to the decrease of urban emissions in the basin during this period. The antipollution measures combined with decreased industrial/transportation activity during the financial crisis are the main factors that are driving a decrease in local urban aerosol emissions. This trend is within the range reported for Europe in the last decade [23].

Meanwhile, the Ångström exponent at 440–870 nm practically shows no trend. SSA at 440 nm decreases at 0.4% per year, which is important compared to the actual range of SSA in the atmosphere. This could be interpreted as a 4% decrease in the examined decade, signifying an increase of absorbing aerosol types. This behavior, combined with the findings of the previous sections of this study, gives some evidence for an increase of absorbing particles, without changing the fine/coarse fraction. This could be explained by a substitution of fine particles with more absorbance at 440 nm, such as brown carbon [75].

![Graphs showing trends of AOD, Ångström exponent, and SSA](image)

**Figure 13.** Anomalies of AOD, Ångström exponent and SSA and deseasonalized trends using AERONET retrievals.

Table 2 presents the deseasonalized total and seasonal trends for the period, as calculated by AERONET, CAMS and MODIS-Terra and MODIS-Aqua. All months with at least 15 days of data were used to calculate the monthly deseasonalized trends from AERONET data. July is the only month that fulfilled this criterion in all years, while other months have gaps in the range of 1–3 years. MODIS-Terra, MODIS-Aqua and CAMS datasets have full datasets. November has a huge decrease since November 2011, when the AOD mean value is 0.28 (AERONET), which is probably attributed to eastern European wildfires. The peak is also detected by all datasets. All datasets used reported a decrease of AOD in the period. Differences could be partially explained by different wavelengths and the gaps in the AERONET database. When collocated data are used (with only the months available for AERONET, used in all datasets), the MODIS-Aqua trend is −1.4%, while the trend for MODIS-Terra
is −1.5\% and for CAMS is −2.0\%. All annual trends are statistically significant at a confidence interval of 95\%.

Seasonal trends calculated from CAMS are significantly different from the other datasets—in particular for winter period (DJF), when the highest decrease is recorded. Satellite and AERONET trends show the highest decreases during summer, when CAMS estimates the most moderate trend. Ground-based and satellite trends for winter have no statistically significant trend.

### Table 2. Deseasonalized trends annually and per season for AOD 440 nm. AERONET LV2 and CAMS 500 nm, MODIS-TERRA and MODIS AQUA 550 nm.

<table>
<thead>
<tr>
<th>AOD Trends (%) per Year</th>
<th>AERONET 440 nm</th>
<th>CAMS 500 nm</th>
<th>MODIS-TERRA 550 nm</th>
<th>MODIS-AQUA 550 nm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual</td>
<td>−1.1 ± 0.7</td>
<td>−2.5 ± 1.1</td>
<td>−1.7 ± 1.0</td>
<td>−1.7 ± 1.0</td>
</tr>
<tr>
<td>DJF</td>
<td>−0.1 ± 0.9</td>
<td>−4.2 ± 1.5</td>
<td>−1.3 ± 1.4</td>
<td>−1.3 ± 1.4</td>
</tr>
<tr>
<td>MAM</td>
<td>−1.8 ± 0.9</td>
<td>−0.5 ± 1.3</td>
<td>−1.7 ± 1.2</td>
<td>−1.2 ± 1.3</td>
</tr>
<tr>
<td>JJA</td>
<td>−3.2 ± 1.1</td>
<td>−1.9 ± 1.5</td>
<td>−2.9 ± 1.3</td>
<td>−2.7 ± 1.4</td>
</tr>
<tr>
<td>SON</td>
<td>−0.8 ± 1.0</td>
<td>−2.1 ± 1.4</td>
<td>−2.6 ± 1.3</td>
<td>−2.6 ± 1.3</td>
</tr>
</tbody>
</table>

### 4. Conclusions

A decade of AERONET measurements have been studied for the city of Athens, Greece, for the 2008–2018 period. Version 3 Level 2.0 data have been used. Maximum AOD monthly values for 340–440 nm were recorded during August and for 675–1640 nm during May. CAMS data also show a maximum at both 550 and 1240 nm during May and a similar seasonal pattern to AERONET retrievals. The mean monthly Ångström exponent at 440–870 nm had maximum values in July and lowest values in April, which was attributed to anthropogenic sources (including biomass burning) and Sahara dust intrusion, respectively. The mean monthly size distribution reveals a maximum of coarse mode in May and a maximum of fine mode in August. SSA also has lowest values in December–January. All the findings lead to the conclusion that the main seasonal pattern in the area is a maximum AOD during summer with a dominance of fine-mode aerosols and frequent dust events in spring time. Additionally, the more absorbent types of aerosols are found during winter.

PM\textsubscript{10} exhibits higher concentrations during winter months and lower concentrations during summer. AOD and PM\textsubscript{10} are more frequently correlated when PBLH is higher than 1200 m. Higher PM\textsubscript{10} concentrations are related with coarse aerosols in the columnar mixture, while more absorbing aerosol types are linked with PM\textsubscript{10} concentrations in the range of 40–60 µg/m\textsuperscript{3}.

Polluted and anthropogenic–mixed are the aerosol classes which are most frequently dominant in the area, comprising 50\% of the cases. Dust and biomass burning aerosols constitute 21\% of the cases when higher AOD values are attributed to these types. Additionally, dust and biomass burning aerosol dominance is more frequently related to PM\textsubscript{10} concentrations higher than 50 µg/m\textsuperscript{3}.

CAMS aerosol types AOD show a constant sulfate aerosol background, with higher dust aerosols in spring time and maximum organic matter in August. The Dust AOD seasonal pattern in CAMS data drives the seasonal pattern of AOD at 1240 nm.

AOD data retrieved from AERONET show a decreasing trend of 1.1\% per year for the area, which could be attributed to antipollution measures and the financial crisis. Satellite and CAMS datasets also show a decrease for the period. The Ångström exponent shows practically no trend. SSA exhibits a 0.4\% decrease per year, which signifies the increase of absorbing aerosol types in the mixture. Satellite and AERONET datasets show the highest decrease during summer months. CAMS seasonal trends have a different pattern than the other datasets.

The climatological results presented in this study are only an early step towards understanding the behavior of aerosols in the complex environment of a European city; longer records are needed to confirm the recorded trends and fully characterize the seasonal patterns. Further analysis combining more sources of data could more clearly provide the signatures of aerosol types and their stratification.
Additionally, similar studies should be performed in other areas in the Eastern Mediterranean area to generalize the findings.

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

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