

Solar absorption by elemental and brown carbon determined from spectral observations

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Black carbon (BC) is functionally defined as the absorbing component of atmospheric total carbonaceous aerosols (TC) and is typically dominated by soot-like elemental carbon (EC). However, organic carbon (OC) has also been shown to absorb strongly at visible to UV wavelengths and the absorbing organics are referred to as brown carbon (BrC), which is typically not represented in climate models. We propose an observationally based analytical method for rigorously partitioning measured absorption aerosol optical depths (AAOD) and single scattering albedo (SSA) among EC and BrC, using multiwavelength measurements of total (EC, OC, and dust) absorption. EC is found to be strongly absorbing (SSA of 0.38) whereas the BrC SSA varies globally between 0.77 and 0.85. The method is applied to the California region. We find TC (EC + BrC) contributes 81% of the total absorption at 675 nm and 84% at 440 nm. The BrC absorption at 440 nm is about 40% of the EC, whereas at 675 nm it is less than 10% of EC. We find an enhanced absorption due to OC in the summer months and in southern California (related to forest fires and secondary OC). The fractions and trends are broadly consistent with aerosol chemical-transport models as well as with regional emission inventories, implying that we have obtained a representative estimate for BrC absorption. The results demonstrate that current climate models that treat OC as nonabsorbing are underestimating the total warming effect of carbonaceous aerosols by neglecting part of the atmospheric heating, particularly over biomass-burning regions that emit BrC.

short lived climate pollutants | aerosol forcing

Black carbon (BC) emitted from combustion sources such as automobile exhaust and biomass burning (1–3) absorbs solar radiation in both the visible and the near-infrared spectra and is estimated to be a principal contributor to global atmospheric warming (4). The short atmospheric lifetime of BC aerosol particles, typically of the order of 1 wk (5, 6), compared with greenhouse gases (which have atmospheric lifetimes of several years or decades) results in BC being not well mixed in the atmosphere but instead geographically and temporally correlated to emission sources. For this reason, reducing BC emissions is an attractive control strategy for climate change that is expected to have a more immediate and regional impact (4, 7, 8). The state of California appears to be a successful example where aggressive control policies for vehicular diesel emissions and domestic wood burning have produced a near 50% decrease in BC concentrations (9). This decline in conjunction with the near-static concentrations of primarily scattering aerosol particles (such as sulfates) may have led to a large negative change in the direct radiative forcing (9).

A simplification in such model estimates of aerosol forcing is that BC is considered to be equivalent to elemental carbon (EC), and the organic fraction of carbonaceous aerosols [organic carbon (OC)] is treated as scattering and is therefore found to have a cooling impact on the climate (10, 11). It is important to realize not only that BC from all emission sources contains both elemental and organic fractions (12), but also that nonsoot OC, particularly that emitted from biomass burning processes, has a significant absorbing component at short wavelengths that may be comparable to the EC absorption (13–17). Consequently, OC that is weakly absorbing in the visible and near-UV spectra may

also contribute to the warming potential of aerosols. In addition to their differing spectral dependence, the wide range of optical properties reported for EC, OC, and mineral dust in the literature (18–22) leads to uncertainties in estimating the absorption attributable to these species. Whereas the attribution of ambient aerosol absorption to EC may be a reasonable approximation in areas dominated by fresh soot emissions, it may lead to misleading estimates of the aerosol forcing when other light-absorbing particles are present.

In more complex environments, a separation of the total absorption into different chemical species is therefore essential, both for constraining the large uncertainties in current aerosol-forcing estimates (23) and for informing any emissions-based control policy. Recent studies reporting the fractions of EC, OC, and mineral dust absorption follow one of three general approaches. First, measured optical and size distribution properties of the aerosol are related to spatial emission patterns to determine the dominant absorbing species (24–27); second, detailed aerosol transport and chemistry models are used in conjunction with assumed optical properties to estimate absorption that is validated against measurements (28–33); and third, closure studies are performed to relate collocated chemically resolved measurements of aerosol mass with optical measurements (34, 35). A recent study by Chung et al. (36) exploited the wavelength dependence of the absorption aerosol optical depths (AAOD) of BC, brown carbon (BrC), and dust aerosols to resolve their relative contribution to the observed absorption optical depths, but used published values for the wavelength dependence of AAOD for OC (brown carbon) and assumed the single scattering albedo (SSA) for EC and OC instead of self-consistently determining these critical properties from observations. This study uses the formalism of Chung et al. without invoking the assumptions noted above. Specifically, the present approach offers the following major improvements: (i) By solving for BC and BrC simultaneously we eliminate the need for assumption of total carbonaceous aerosol (TC) properties that depend on source-specific compositions. (ii) The OC AAE is determined empirically by exploiting measurements at the longer wavelengths of 870 and 1020 nm. (iii) We consider scattering optical depths to better constrain the phase space on the basis of size. (iv) Finally, we also offer empirical estimates of the SSA. The proposed empirical scheme requires only a single set of aerosol optical measurements, allowing for spatial and temporal coverage on a nearly global scale. We provide a robust set of constraints that will allow climate modelers to improve the estimation of the BrC aerosol forcing.

The remainder of this paper is organized as follows: We describe the equations composing our partitioning scheme; the

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equations are parameterized using an observational dataset as input; the scheme uses observations from California as a test case to determine EC, OC, and dust absorption; our estimates are compared with model predictions and emission inventories; and finally we comment on the limitations of our scheme and the scope for future applications.

Partitioning AAOD

The total aerosol optical depth (AOD) represents the total light extinction due to scattering and absorption by an aerosol and is related to the AAOD through the SSA as

$$\text{AAOD}(\lambda) = (1 - \text{SSA}(\lambda)) \times \text{AOD}(\lambda). \quad [1]$$

The AOD and AAOD are extensive properties that depend on the total aerosol concentration, whereas the SSA (the fraction of total aerosol extinction due to scattering) is an intensive property that depends only on the aerosol composition; however, the value of all three depends on the wavelength of the incident light. The spectral dependence of the AOD and AAOD is related to reference values with an exponential dependence

$$\text{AOD}(\lambda) = \text{AOD}_{\text{ref}} (\lambda/\lambda_{\text{ref}})^{-\text{EAE}} \quad [1]$$

$$\text{AAOD}(\lambda) = \text{AAOD}_{\text{ref}} (\lambda/\lambda_{\text{ref}})^{-\text{AAE}}. \quad [2]$$

In the classical formulation, the extinction angstrom exponent (EAE) and absorption angstrom exponent (AAE) are thought to be constants; however, recent studies have shown them to depend on the wavelength, particle size, and particle composition (37–39). Because the total AAOD is an extensive property, the contributions due to EC, OC, and dust are additive such that

$$\text{AAOD}(\lambda) = \text{AAOD}_{\text{EC}}(\lambda) + \text{AAOD}_{\text{OC}}(\lambda) + \text{AAOD}_{\text{dust}}(\lambda). \quad [3]$$

Eqs. 2 and 3 can be combined to generate a set of independent equations that compose our primary formulation:

$$\begin{aligned} \text{AAOD}(\lambda_i) = & \text{AAOD}_{\text{ref,EC}} (\lambda_i/\lambda_{\text{ref}})^{-\text{AAE}_{\text{EC},i}} \\ & + \text{AAOD}_{\text{ref,OC}} (\lambda_i/\lambda_{\text{ref}})^{-\text{AAE}_{\text{OC},i}} \\ & + \text{AAOD}_{\text{ref,dust}} (\lambda_i/\lambda_{\text{ref}})^{-\text{AAE}_{\text{dust},i}}. \end{aligned} \quad [4]$$

If the AAOD is known in at least three wavelengths (i.e., $i = 1, 2, 3$), the three-part Eq. 4 can be solved for the three unknown reference AAOD values corresponding to EC, OC, and dust, which can then in turn be used to partition the AAOD at any wavelength (Fig. S4). The complete set of AAE values parameterizes these equations and constrains the phase space in which physically relevant solutions exist. There are only two inherent assumptions in this formulation: First, the AAE values for the component absorbing species EC, OC, and dust are intrinsic properties that are not dependent on the mixing state; and second, the measured AAOD value represents a well-mixed sample of these species.

The full implementation of this method requires knowledge of the AAE values to partition the AAOD and of the SSA values to determine the AOD. In theory a number of pathways can be used to determine these parameters, but as stated in the introductory section, in this work we rely only on empirical values derived from optical measurements of ambient aerosols. Finally, because the absorption and scattering components of extinction can be related to aerosol chemistry and size, respectively (40), we use the scattering angstrom exponent (SAE) in place of the EAE for the remainder of this work.

The ground-based Aerosol Robotic Network (AERONET), a globally distributed network of automated sun and sky radiometers (41), provides long-term, continuous, and readily available measurements of aerosol optical properties that are an ideal resource for this study. In addition to multiband retrievals of AOD between 340 and 1,020 nm from direct sun measurements, the inversion algorithm provides SSA (and corresponding AAOD) estimates from sky radiance measurements at 440, 675, 870, and 1020 nm (42).

Absorption Angstrom Exponent

Because the primary focus of this work is partitioning absorption in the visible and near-UV spectra, we select the three shortest AERONET wavelengths for the remainder of our analysis. We designate the angstrom exponents for AOD and AAOD between 440 and 675 nm as SAE1 and AAE1 and similarly the angstrom exponents between 675 and 870 nm as SAE2 and AAE2, respectively.

Several AERONET sites are dominated by emissions corresponding to a single absorbing aerosol (25). To determine the angstrom exponents and SSA for the pure species in Eq. 4, we have selected sites typically dominated by specific sources related to dust, EC, and OC with details in Table S1. In addition to providing

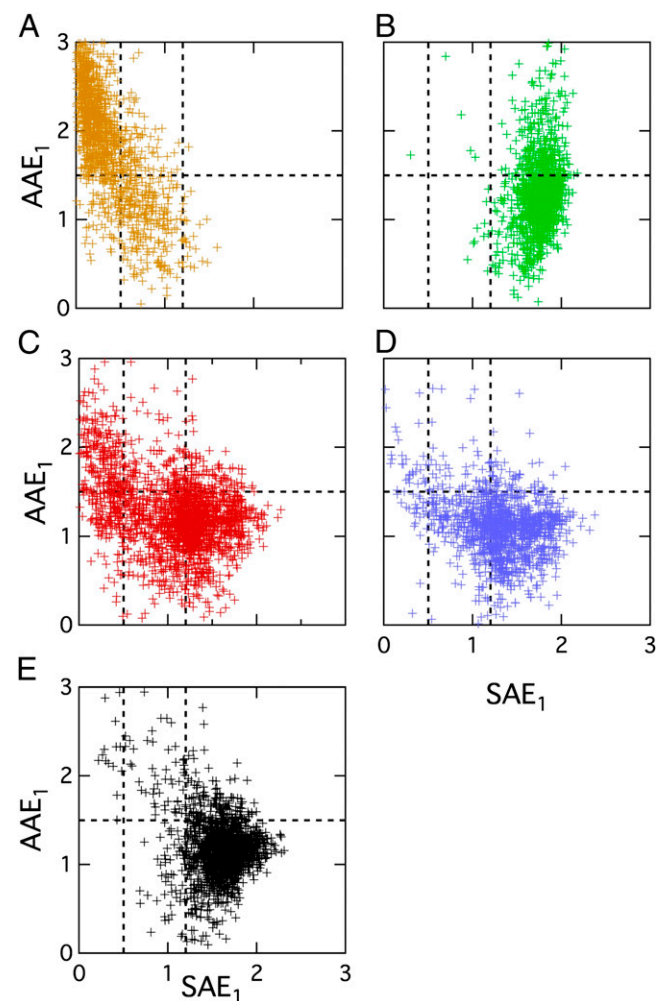


Fig. 1. (A–E) Scatter plots of AAE1 and SAE1 calculated from AERONET measurements at (A) dust (DU), (B) BB, (C) UF, (D) NF, and (E) CA sites. Dashed lines illustrate the threshold values of SAE1 = 0.5, SAE1 = 1.2, and AAE1 = 1.5 used to separate dust-dominated and dust-free regimes.

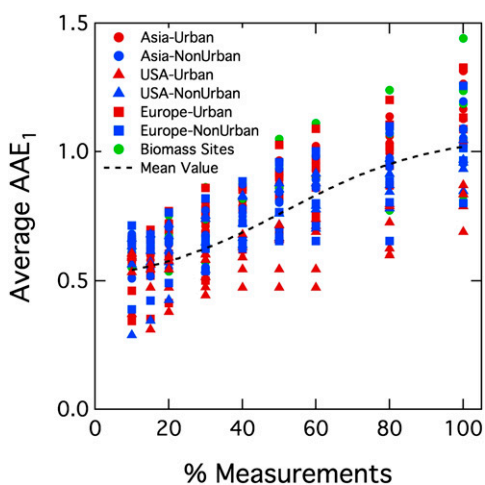


Fig. 2. Mean value of AAE1 determined by averaging the frequency distribution for AERONET sites in Table S1. The dashed line shows a fit to the mean value.

insights into pure component properties, the varying geographic location and sources associated with these sites allow us to test our hypothesis that AAE and SSA are intrinsic properties.

Dust aerosols have a crustal origin distinct from the combustion-related origin of carbonaceous aerosols and have been shown to have significantly different wavelength dependences of absorption and scattering (25), allowing us to establish thresholds in AAE1 and SAE1 for identifying dust-dominated and dust-free measurements. We provide the detailed analysis and corresponding figures in *SI Text*. Measurements with SAE1 < 0.5 are found to be dust dominated and for pure mineral dust we find $AAE1 = 2.20 \pm 0.50$ and $AAE2 = 1.15 \pm 0.40$. The conditions $SAE1 > 1.2$ or $AAE2/AAE1 > 0.8$ correspond to dust-free (alternatively, carbon-dominated) measurements (Fig. S1) and AAE1 calculated for total carbon is found to be 1.06 [nonurban fossil fuel (NF)], 1.14 [urban fossil fuel (UF)], and 1.28 [biomass burning (BB)], agreeing well with the value of 1.0 typically reported for BC (20). It is important to note two subtle yet important conclusions based on these values. First, the similarity in magnitude across varied geographical regions and emission sources (Figs. S3 and S4) supports the hypothesis that the absorption angstrom exponent is an inherent intrinsic property. Second, the differences can be attributed to different relative amounts of elemental and organic carbon (43), with biomass-burning aerosols containing a larger organic fraction and therefore having a larger absorption angstrom exponent than fossil fuel aerosols.

Fig. 1 illustrates the relationship between AAE1 and SAE1 for all AERONET measurements included in this study. Whereas the separation between dust-dominated and dust-free measurements based on the threshold values is clear, all fossil fuel and biomass-burning measurements occupy a continuous region of the phase space and the separation of pure EC and pure OC contributions requires additional analysis. Because EC has a weaker spectral dependence than OC, we can posit that the lower end of the AAE1 values reflects absorption due to EC. Fig. 2 illustrates the mean values of AAE1 calculated by averaging different fractions of the total frequency distribution at the different AERONET sites. We find that the mean AAE1 value asymptotically converges to 0.55 ± 0.24 , representing the average low-end baseline for the absorption angstrom exponent. We relate this value to pure EC, again noting that varying amounts of co-occurring OC raise this value in ambient measurements. A similar analysis gives us a value of $AAE2 = 0.83 \pm 0.40$ for EC. These values are found to be independent of the location, the

source, and the total aerosol concentration and therefore can be considered robust representations of pure EC. Because both OC and dust have a stronger spectral dependence than EC, any observed values of AAE lower than these mean values correspond to BC-dominated aerosols, leading to our third constraint: If $AAE1 < 0.55$ and $AAE2 < 0.83$, $AAOD = AAODEC$.

The calculation of OC absorption angstrom exponents requires one additional assumption. Because OC absorbs primarily in the UV wavelength and at shorter visible ($\lambda < 700$ nm) wavelengths (13, 44), we modify Eq. 4 such that

$$AAOD_{OC}(870) = 0. \quad [5]$$

In dust-free regions the total AAOD at 870 nm is entirely due to EC, and the EC AAODs at 675 and 440 nm are calculated using Eq. 2. Finally, Eq. 3 is used to calculate the OC AAODs at these two wavelengths, and the value of AAE1 for OC can be determined (because we assume OC does not absorb at 870 nm, AAE2 is undefined). The frequency distribution of OC AAE1 determined in this fashion is illustrated in Fig. 3 and is found to have a narrow distribution centered around 4.5 for all sites, further supporting our hypotheses that the absorption angstrom exponent is an intrinsic property. The mean value is found to be 4.55 ± 2.01 .

Table 1 summarizes the values of absorption angstrom exponents determined in this work and compares them with the ranges of values reported in the literature, both based on measurements and determined from models using an assumed refractive index. The AAE values for dust and total carbon lie within the ranges reported in the literature and typically agree well with the central values. The AAE values for EC determined in this study are lower than literature values, and the values for OC lie toward the upper end reported in the literature. This indicates that EC and OC (treated as pure chemical species in this study) are typically comingled in combustion studies, even close to the source. Recent modeling studies have examined the effect of internal mixing between EC and OC on the AAE using a Mie-scattering algorithm in conjunction with a core-shell assumption (38, 45). OC coatings (representing internal mixing) are found to change the AAE for BC from 0.7 (no particles coated) to 0.1 (all particles coated), with this range being consistent with the value of 0.55 ± 0.24 calculated in this work. One final caveat needs to be considered in our approach: Any absorption by fine-mode dust (high

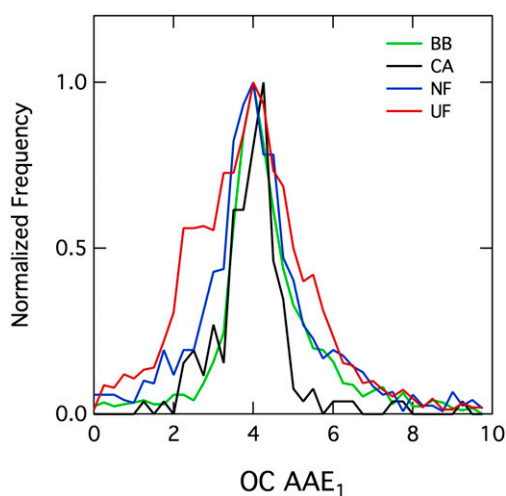


Fig. 3. Normalized frequency distribution of the OC absorption angstrom exponent determined from biomass burning (BB), California (CA), nonurban fossil fuel (NF), and urban fossil fuel (UF) sites between 440 and 675 nm.

Table 1. Absorption angstrom exponents for dust, total carbon, BC, and OC determined in this study and the ranges reported in the literature

Species	This work		Literature, measured		Literature, modeled	
	AAE1	AAE2	AAE	Refs.	AAE	Refs.
Dust	2.20 ± 0.50	1.15 ± 0.50	1.0–4.0	(22, 25, 50, 51)		
Total carbon	1.17 ± 0.40	1.23 ± 0.40	1.0–1.3	(19, 25, 52, 53)	1.0–1.7	(38)
BC/EC/soot	0.55 ± 0.24	0.85 ± 0.40	0.8–1.1	(35, 43)	1.00	(38)
OC	4.55 ± 2.01	—	1.8–7.0	(13, 19, 25, 35, 43, 54–56)	3.0–6.0	(28, 57)

AAE, high SAE) would be masked by absorption due to OC. Fig. 1A illustrates, however, that less than 1% of observations in regions with significant dust absorption lie in this phase space, consistent with most mineral dust being found in the coarse mode (46), and similarly less than 10% of points in the other regions fall in the coarse-dust region. We therefore do not expect significant errors due to fine-dust absorption.

Single Scattering Albedo

Whereas the total AOD can be attributed to BC, OC, and dust using Eq. 4, a similar partitioning of the AOD is not possible due to the variable contribution of purely scattering aerosols. To reconstruct the AOD for the absorbing species, we estimate the pure species single scattering albedo by a recursive numerical approach. The SSA is calculated at sites where each absorbing species is expected to be dominant; for example, BC SSA is determined only from fossil fuel-dominated sites. In summary, the derived SSAs for dust and OC (Fig. S2) are consistent with the range of values reported in the literature (Table S2) and both become more absorbing at shorter wavelengths. The spectral dependence of EC SSA on the other hand is weak and within the error at each bar at each wavelength; therefore, we use the value at 870 nm (with no OC and minimal dust absorption) for EC. We find a higher SSA for EC (0.38) than that typically reported by measurements (0.15–0.30). The published estimates are for aerosol particles generated in the laboratory or calculated in models (GOCART) whereas our SSA is based on ambient observations that are likely to reflect atmospherically processed EC that may have contamination from internally and/or externally mixed organics. This mixing will tend to enhance the aerosol SSA and therefore our SSA values should be considered as an upper limit. Our analysis constrains the EC SSA at 440 nm to values between 0.15 and 0.48, with an average value of 0.38. It should be noted that the derived SSA does not influence the absorption estimates because AAODs were derived independent of the SSAs.

California Case Study

With the complete set of AAE and SSA values, Eq. 4 with its associated constraints for dust-dominated, dust-free, and EC-dominated regimes can be applied to any set of multiwavelength absorption measurements. Here we apply this method to California, which provides an ideal test case due to varied aerosol emissions sources and readily available aerosol measurements and emission inventories. Although the CA AERONET dataset with valid SSA measurements is of insufficient size to construct a complete climatology, we can assess the seasonal and regional trends in aerosol properties and compare them to Goddard Chemistry, Aerosol, Radiation, and Transport (GOCART) model (29) predictions and regional emission estimates (47). The sensitivity of these results to the selected AAE parameterization is examined in Table S2. We define north-central California to be one region (north of 35°N) and southern California to be the other, such that both contain five operational AERONET sites, and the year is divided into two seasons: winter–spring from December to May and summer–fall from June to November. Table 2 summarizes the trends in AOD and AAOD for the absorbing species on the basis of the CA measurements and the GOCART model. Because SSA retrievals are valid only when the total AOD > 0.4 (42), comparing the absolute values results in a significant bias. We therefore report the AOD and AAOD fractions normalized by the total. In the seasonal comparison, we find that dust contributes a higher fraction of the AAOD in the winter–spring months (19% in northern California and 72% in southern California) compared with the summer–fall months (7% in northern California and 3% in southern California). Although the southern winter value may be anomalously high due to a few outliers, this general trend is consistent with trans-Pacific transport of dust seen in California in the winter months (48). We find a much higher absorption attributed to OC in the summer–fall months (27% in the north and 39% in the south) compared with the winter–spring months (15% in the north and 9% in the south), which is consistent with the California wildfires

Table 2. Dust, OC, and BC fraction of the total AAOD and AOD as determined from AERONET (this work) and the GOCART model in California

	Season	<i>n</i>	This work 440 nm			GOCART 550 nm			This work 675 nm		
			Dust	OC	BC	Dust	OC	BC	Dust	OC	BC
AAOD											
Northern and central California (>35°N)	December–May	77	0.19	0.15	0.66	0.51	0.11	0.38	0.12	0.03	0.84
	June–November	150	0.07	0.27	0.65	0.46	0.11	0.43	0.05	0.07	0.88
Southern California (<35°N)	December–May	35	0.72	0.09	0.18	0.50	0.10	0.40	0.64	0.03	0.33
	June–November	66	0.03	0.39	0.57	0.38	0.11	0.52	0.02	0.11	0.87
AOD											
Northern and central California (>35°N)	December–May	26	0.11	0.05	0.08	0.25	0.05	0.02	0.04	0.01	0.06
	June–November	87	0.05	0.11	0.10	0.23	0.05	0.02	0.02	0.01	0.09
Southern California (<35°N)	December–May	35	0.52	0.03	0.03	0.25	0.05	0.02	0.36	0.01	0.05
	June–November	66	0.01	0.14	0.07	0.19	0.05	0.03	0.01	0.03	0.12

associated with these months (49). BC makes up between 57% and 60% of the AAO in all cases (Fig. S6), indicating that there are no seasonal or regional trends in California fossil fuel emissions. Regionally, dust comprises a higher fraction of the AAO in northern California (consistent with transport events) and OC comprises a higher fraction in the south. The regional pattern may be an indication of absorption due to secondary organics that are typically present in high concentrations in the urban areas of southern California. We find good agreement in the OC fraction between our observationally constrained method and the GOCART predictions, with calculated values between 15% and 39% at 440 nm and 3% and 11% at 675 nm, bracketing the modeled fraction of 11% at 550 nm. We find a lower contribution and higher variability in the dust (7–72%) compared with those of GOCART, which attributes between 38% and 51% of AAO to dust. One possible explanation for this difference is that the AERONET sites are primarily located along the coast and may not be sampling dust lofted from exposed lakebeds in inland California and the desert regions of Nevada and Arizona. Finally, we find that the three absorbing species compose between 16% and 52% of the total AOD, which is typically lower (except the anomalously high southern California winter value) than the 27–32% predicted by GOCART, indicating that the aerosol represented by AERONET measurements is more scattering. This difference can again be explained by the influence of the coastal sites that typically have a larger fraction of optical extinction due to purely scattering sea-salt and sulfate particles. The total AOD and AAO (at 675 nm) for dust, EC, and OC are illustrated in Fig. 4A and B, respectively: Although these are based on high pollution events, they still agree with the regional trends in emissions illustrated in Fig. 4C, indicating that our method provides a reasonable partitioning between the three species. Finally, Fig. 4D provides a unique wavelength-dependent

perspective on the relative contribution of OC absorption: Although it is negligible at long wavelengths, it comprises up to 50% of the EC absorption at short wavelengths, indicating that the treatment of OC as being absorbing is critical for successful estimations of aerosol forcing. This result is consistent with measurements conducted in the Los Angeles basin that attributed between 25% and 30% of UV light absorption to organic carbon species and between 50% and 62% to black carbon particulates (17).

Conclusions

We have proposed an empirical method for determining the fraction of aerosol absorption attributable to dust, OC, and EC that exploits the differences in wavelength dependence for the principal absorbing species. We have self-consistently determined the AAE and SSA values associated with these species on the basis of a large global ensemble of AERONET measurements dominated by specific emission types. The parameter values are based entirely on observations and therefore do not require the assumption of aerosol chemical and physical properties such as refractive index, mixing state, or size distributions. These values are found to be consistent with those reported in the literature and in use in current aerosol models, in particular the AAE value for total carbon close to the theoretical value of 1.0. The method has been applied to California as a test case, where we find an enhanced absorption due to dust in the winter months and due to OC in the summer months. We find that OC on average contributes 28% to the total absorption at the shortest wavelength (440 nm), providing a good first-order estimate for the so-called brown carbon absorption that is presently poorly quantified.

We conclude by commenting on the general applicability of this method. Although we recommend values for the model

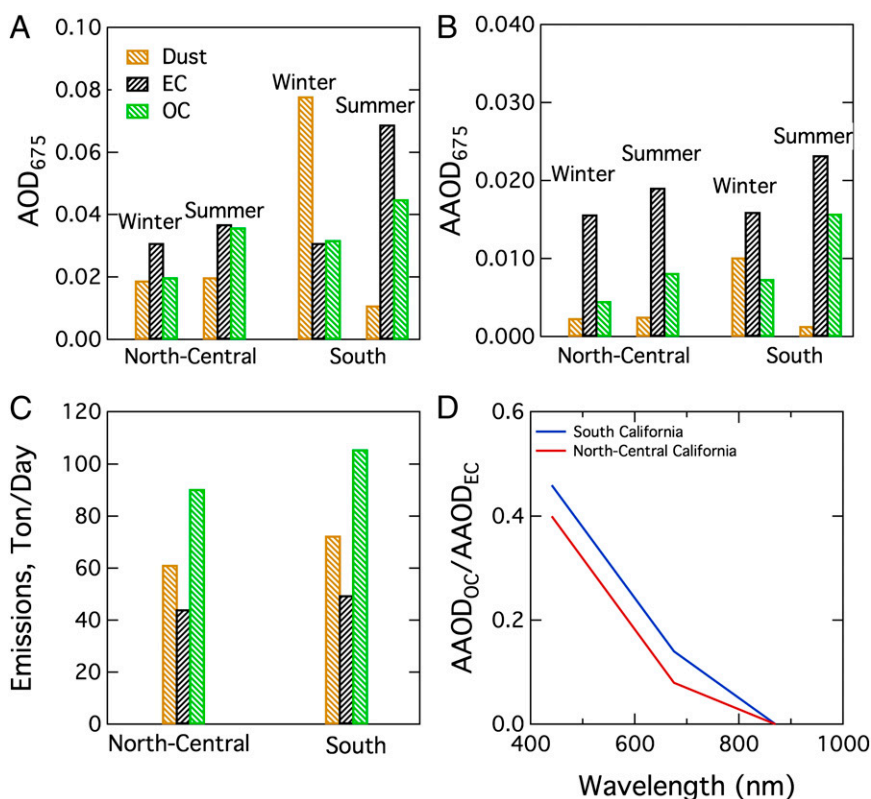


Fig. 4. (A–C) Species-resolved (A) AOD, (B) AAO, and (C) emissions in California for EC, OC, and dust. (D) Wavelength-dependent relative absorption of OC and EC.

parameters based upon a global dataset, the equations can be fine tuned to any region by following the procedures outlined here for determining the SSA and AAE values. Because the only required input for the equation is a multiband measurement of the total absorption depth, the AERONET results reported here can be extended to the large ensembles of available diverse

laboratory, ground-based, and remote satellite measurements. Finally, the analyses indicate the existence of significant BrC absorption that has been largely overlooked in climate models.

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