

Aerosol Effects on Clear-Sky Shortwave Heating in the Asian Monsoon Tropopause Layer

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Key Points:

- The Asian tropopause aerosol layer produces a 5–25% direct enhancement of clear-sky shortwave heating above the summer monsoon
- Effects are largest where shortwave heating is weakest, with similar magnitudes to water vapor and ozone effects near the monsoon tropopause
- Discrepancies across recent aerosol analysis and forecast products cause large uncertainties in aerosol forcing of heating and fluxes

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Abstract

The Asian Tropopause Aerosol Layer (ATAL) has emerged in recent decades to play a prominent role in the upper troposphere and lower stratosphere above the Asian monsoon. Although ATAL effects on surface and top-of-atmosphere radiation budgets are well established, the magnitude and variability of ATAL effects on radiative transfer within the tropopause layer remain poorly constrained. Here, we investigate the impacts of various aerosol types and layer structures on clear-sky shortwave radiative heating in the Asian monsoon tropopause layer using reanalysis products and offline radiative transfer simulations. ATAL effects on shortwave radiative heating based on the MERRA-2 aerosol reanalysis are on the order of 10% of mean clear-sky radiative heating within the tropopause layer, although discrepancies among recent reanalysis and forecast products suggest that this ratio could be as small as $\sim 5\%$ or as large as $\sim 25\%$. Uncertainties in surface and top-of-atmosphere flux effects are also large, with values spanning one order of magnitude at the top-of-atmosphere. ATAL effects on radiative heating peak between 150 hPa and 80 hPa (360 K–400 K potential temperature) along the southern flank of the anticyclone. Clear-sky and all-sky shortwave heating are at local minima in this vertical range, which is situated between the positive influences of monsoon-enhanced water vapor and the negative influence of the ‘ozone valley’ in the monsoon lower stratosphere. ATAL effects also extend further toward the west, where diabatic vertical velocities remain upward despite descent in pressure coordinates.

Plain Language Summary

Every summer, a layer of polluted air laden with aerosol particles collects above the convective storms of the Asian monsoon as part of a broad upper-level circulation centered over the Tibetan Plateau. Researchers have developed a working understanding of how the dynamical environment shapes this Asian tropopause aerosol layer. The motivating question for this work is: how might the aerosol layer reshape its environment? Aerosols can absorb and scatter sunlight, affecting both the amount of sunlight transmitted through the layer and the magnitude of solar heating within the layer. These effects depend on aerosol species and their vertical distribution within the layer, both of which are highly variable. In this paper, we translate variations and uncertainties in the amount, composition, and vertical distribution of aerosols near the Asian monsoon tropopause into variations and uncertainties in the absorption and scattering of solar radiation by the aerosol layer. We find that aerosols account for a substantial part (5-25%) of heating by solar radiation near the tropopause. The vertical location and horizontal extent of the aerosol effects are distinct from those of other radiative effects.

1 Introduction

Aerosol effects on radiative heating are among the most uncertain and complex challenges to understanding atmospheric variability at scales ranging from the development of weather systems to the long-term evolution of climate (Ramaswamy et al., 2018; Szopa et al., 2021). Different aerosol types have different direct effects on atmospheric radiative transfer. Direct effects, also referred to as radiative forcing from aerosol–radiation interactions, can be broadly classified into absorption and scattering by atmospheric aerosols. For example, black carbon (BC), brown carbon (a form of organic carbon; OC), and some mineral dust particles can directly absorb solar radiation (K.-M. Lau & Kim, 2006; Samset et al., 2018), while sulfate, nitrate, and most dust and OC aerosols act mainly to scatter solar radiation (Whitby, 1978; Kinne et al., 2006). In addition to these direct effects, indirect radiative effects include, for example, aerosol-induced changes in the occurrence frequencies, lifetimes, and optical properties of clouds (Twomey, 1977; Albrecht, 1989; Lohmann, 2017; Kreidenweis et al., 2019). As a general rule, and with the exception of very large aerosol particles, direct forcing by aerosol–radiation interactions mainly in-

64 influences the shortwave (solar) part of the spectrum, while indirect effects influence both
65 longwave (thermal) and shortwave radiative transfer (Charlson et al., 1992).

66 Aerosol distributions are highly variable in space and time owing to spatial gra-
67 dients in emissions, secondary formation processes, and spatiotemporal heterogeneity in
68 the efficiency of removal processes (Kreidenweis et al., 2019). The prevalence of inter-
69 nal mixing and large uncertainties in aerosol optical properties further complicate the
70 identification of aerosol types and their impacts on atmospheric radiation. As a conse-
71 quence, there remain substantial uncertainties in the magnitude of direct aerosol radia-
72 tive forcings at the surface and top-of-atmosphere (Ramaswamy et al., 2018; Kuniyal &
73 Guleria, 2018; Szopa et al., 2021), let alone in the vertical profile of radiative heating.
74 These challenges are especially acute with respect to the structure and radiative effects
75 of the Asian tropopause aerosol layer (ATAL; Vernier et al., 2011), which develops each
76 year during boreal summer in the upper troposphere and lower stratosphere (UTLS) above
77 the Asian summer monsoon (Vernier et al., 2015; Yu et al., 2015; Bian et al., 2020).

78 The Asian summer monsoon is among the most important sources of water vapor
79 and tropospheric pollutants to the global stratosphere (Fu et al., 2006; Randel et al., 2010;
80 Pan et al., 2016; Ploeger et al., 2017; Yu et al., 2017; Lelieveld et al., 2018). Strong deep
81 convection associated with the monsoon pumps moist, polluted air (Park et al., 2007;
82 Pan et al., 2016; Bian et al., 2020) upward into a vigorous upper-level anticyclone bounded
83 by the tropical easterly jet to the south and the subtropical westerly jet to the north (Hoskins
84 & Rodwell, 1995; Garny & Randel, 2013; Legras & Bucci, 2020). Much of the air within
85 this upper-level monsoon anticyclone traces back to the boundary layer over South and
86 East Asia (Bergman et al., 2013; Orbe et al., 2015; Vogel et al., 2016; Zhang et al., 2020),
87 where abundant emissions of aerosols and aerosol precursors contribute to ATAL forma-
88 tion (Neely et al., 2014; Yu et al., 2015; Vernier et al., 2017; Bian et al., 2020). Enhanced
89 aerosol concentrations in the UTLS alter radiative heating at the tropopause level both
90 directly (Toohey et al., 2014; Vernier et al., 2015; Yu et al., 2015; Fadnavis et al., 2017,
91 2019) and indirectly through their interactions with clouds (Su et al., 2011; Dong et al.,
92 2019; Fadnavis et al., 2019). However, the practical impacts of these aerosol effects de-
93 pend in large part on the composition and vertical structure of the aerosol layer, which
94 are highly variable (Hanumanthu et al., 2020) and poorly constrained (Bian et al., 2020).

95 Estimates of ATAL composition are based mainly on numerical models, some of
96 which indicate that sulfate aerosols originating from East Asia are the leading compo-
97 nent (Neely et al., 2014; Vernier et al., 2015; Yu et al., 2015), while others point to ni-
98 trate aerosols (Gu et al., 2016) and still others point to leading roles for organic aerosols
99 or dust (W. K. M. Lau et al., 2018; Ma et al., 2019; Bossolasco et al., 2021). Observa-
100 tional studies conducted on ATAL composition to date do not support large concentra-
101 tions of dust at the tropopause level (Vernier et al., 2017) and highlight a surprising abun-
102 dance of nitrate aerosols (Höpfner et al., 2019), which are still not represented in many
103 models (Kreidenweis et al., 2019; Bossolasco et al., 2021). Other observational studies,
104 while unable to measure aerosol composition, have emphasized large variations in ATAL
105 amplitude and vertical structure at daily time scales (Brunamonti et al., 2018; Zhang
106 et al., 2020; Hanumanthu et al., 2020; Mahnke et al., 2021). These variations and asso-
107 ciated uncertainties complicate efforts to constrain the radiative effects of the ATAL.

108 The ATAL is distinct from episodic volcanic aerosol loading of the lower stratosphere
109 in its composition, seasonality, vertical location, and longitudinal extent (Bian et al., 2020).
110 Although much progress has been made in evaluating the variability, formation mech-
111 anisms, and tropospheric sources of the ATAL, its effects on the energy budget and ther-
112 modynamic structure of the tropopause layer are not yet clear. Important steps in this
113 direction include the model-based analyses conducted by Yu et al. (2015) and Fadnavis
114 et al. (2017, 2019), along with a recent reanalysis-based dynamical assessment of ATAL
115 evolution as observed by satellite (He et al., 2021). However, given inter-model discrep-
116 ancies and observational uncertainties in the composition and structure of the ATAL,

117 the extent to which the results of these studies can be generalized to other model sys-
118 tems or the natural atmosphere remains unclear.

119 As a step toward addressing this gap, we adopt an idealized framework to exam-
120 ine ATAL effects on radiative heating in the monsoon UTLS, focusing on clear-sky short-
121 wave heating. We focus on shortwave effects for three reasons. First, aerosol radiative
122 forcing are mainly confined to the shortwave part of the spectrum, especially for the re-
123 latively small aerosol particles that predominate within the ATAL (Vernier et al., 2017;
124 Mahnke et al., 2021; Weigel et al., 2021). Second, the shortwave effects of aerosol per-
125 turbations in the UTLS can be treated as a diabatic forcing while longwave responses
126 are often dominated by thermal relaxation (Toohey et al., 2014). Third, convective ven-
127 tilation of the UTLS above the monsoon also creates regional-scale anomalies in ozone
128 and water vapor, both of which play important roles in shortwave radiative transfer near
129 the tropopause. These anomalies provide convenient comparison points for evaluating
130 the relative influences of ATAL on shortwave heating in the UTLS. Further restricting
131 our analysis to clear-sky conditions is motivated mainly by the ATAL being located above
132 the majority of convective anvil clouds (Vernier et al., 2015; Bian et al., 2020). Although
133 aerosols also have important impacts on the frequency, distribution, and optical prop-
134 erties of tropopause-level cirrus clouds (Su et al., 2011; Riuttanen et al., 2016; Fadnavis
135 et al., 2019), these interactions are poorly constrained. Accounting for aerosol effects on
136 cirrus clouds would thus compound already large uncertainties while requiring evalua-
137 tion of aerosol–cloud covariability, and is deferred to future work.

138 This paper is organized in five parts. In Section 2, we introduce the reanalysis and
139 forecast products and the libRadtran radiative transfer model. In Section 3, we exam-
140 ine the spatial and temporal distributions of aerosol in the Asian monsoon UTLS accord-
141 ing to the MERRA-2 reanalysis and describe a series of idealized sensitivity experiments
142 exploring the clear-sky shortwave effects of variations in the amplitude, composition, and
143 vertical structure of the ATAL. In Section 4, we contextualize the ATAL radiative ef-
144 fects by comparing their magnitudes and distributions to those of ozone, water vapor,
145 and clouds, as well as quantifying the impacts of uncertainties in ATAL composition and
146 structure across recent reanalysis and forecast products. In Section 5, we provide a short
147 summary of the conclusions and possible next steps.

148 2 Data and Methods

149 2.1 The MERRA-2 aerosol reanalysis

150 The primary dataset for this study is the Modern-Era Retrospective Analysis for
151 Research and Applications, version 2 (MERRA-2; Gelaro et al., 2017) for the 10-year pe-
152 riod 2011–2020. MERRA-2 is unique among current meteorological reanalyses (see Wright
153 et al., 2022, their Table 2.12) in that it includes an interactive aerosol analysis (Randles
154 et al., 2017). Aerosols are simulated via the Goddard Chemistry, Aerosol, Radiation, and
155 Transport (GOCART) model, which considers sources, sinks, and chemical properties
156 of 15 aerosol types and classes, including dust (five non-interacting size bins), sea salt
157 (five non-interacting size bins), sulfate, hydrophilic and hydrophobic BC, and hydrophilic
158 and hydrophobic OC (Chin et al., 2002; P. Colarco et al., 2010; P. R. Colarco et al., 2014;
159 Randles et al., 2017). Observations used in the data assimilation are limited to remote
160 sensing measurements of aerosol optical depth (AOD) and do not alter the composition
161 or relative vertical distribution of aerosols (Randles et al., 2017).

162 Although prescribed emissions to the aerosol model vary in time over much of the
163 1980–2010 period, most emissions sources use either constant or annually-repeating monthly
164 values over the analysis period 2011–2020 (Randles et al., 2017). This includes volcanic
165 emissions, which only include a repeating annual cycle of outgassing (omitting eruptions)
166 after 2010. Injections of sulfate to the stratosphere by volcanic eruptions can make it dif-

167 difficult to distinguish the ATAL (Thomason & Vernier, 2013; Vernier et al., 2015). Lim-
 168 iting our analysis to 2011–2020 essentially eliminates this potential confounding effect.
 169 Biomass burning emissions are from version 2.4 of the Quick Fire Emissions Database (QFED;
 170 Darnenov & da Silva, 2015). MERRA-2 has been shown to perform well with respect
 171 to independent observations of AOD and fluxes (Randles et al., 2017), as well as absorb-
 172 ing aerosol optical depth, ultraviolet index, and vertical structure (Buchard et al., 2017).
 173 The ATAL in MERRA-2 has previously been examined by W. K. M. Lau et al. (2018),
 174 who focused on carbonaceous aerosols and dust during the pre-monsoon and peak mon-
 175 soon periods (May–August) of 2008. Their results showed that MERRA-2 produces a
 176 well-defined ATAL fed mainly by deep convection over North India and the Sichuan Basin.
 177 Further details on emissions, previous validation, and the rationale for using MERRA-
 178 2 as the basis for this work are provided in Supporting Information (Text S1).

179 MERRA-2 products used in this work include daily-mean vertical profiles of tem-
 180 perature, specific humidity, mass mixing ratios of ozone and aerosol species, pressure,
 181 and geopotential height; temperature tendencies due to clear-sky radiative heating on
 182 model levels for the analysis window 04:30–07:30 UTC (corresponding to mid-day in our
 183 core analysis region); and hourly surface albedo (Table S1 in Supporting Information).
 184 To capture the largest concentrations of aerosol in the ATAL, we focus mainly on mean
 185 distributions and profiles from MERRA-2 for July–September 2011–2020. Daily-mean
 186 data from May–September are used to illustrate the seasonal evolution of the MERRA-
 187 2 ATAL and its effects on clear-sky shortwave heating in the tropopause layer during 2011–
 188 2020, and daily means for July–September are used to assess variability. Regional selec-
 189 tions are used to show variations in ATAL properties and radiative effects as functions
 190 of longitude (50°E–120°E, meridionally averaged over 22.5°N–25°N) and latitude (18°N–
 191 42°N, zonally averaged over 87.5°E–90°E) across the monsoon anticyclone. These two
 192 vertical cross-sections intersect at 22.5°N–25°N and 87.5°E–90°E, north of the Bay of Ben-
 193 gal. This 2.5°×2.5° grid cell is designated the ‘core’ region and is used to define the base-
 194 line for idealized radiative transfer calculations.

195 2.2 Radiative transfer model

196 Offline radiative transfer calculations to quantify ATAL effects on clear-sky short-
 197 wave radiative heating are conducted using version 2.0.3 of the libRadtran radiative trans-
 198 fer model (Mayer & Kylling, 2005; Emde et al., 2016), a multi-layer model developed to
 199 support flexible representation of absorption and scattering in the atmosphere. libRad-
 200 tran is centered around the uvspec radiative transfer code, which can simulate radiative
 201 transfer across the solar and thermal spectra at a range of spectral resolutions (Mayer
 202 & Kylling, 2005; Emde et al., 2016). Default aerosol optical properties are from the Op-
 203 tical Properties of Aerosols and Clouds (OPAC) database (Hess et al., 1998), including
 204 radiative interactions of ten species of aerosols over wavelengths ranging from 250 nm
 205 to 40 μm across eight grades of relative humidity. OPAC also serves as the basis for aerosol
 206 optical properties in MERRA-2 (Chin et al., 2002; Randles et al., 2017; Buchard et al.,
 207 2017). Applications of libRadtran to shortwave radiative transfer and aerosol effects near
 208 the tropopause include development of remote sensing retrieval algorithms (Theys et al.,
 209 2007; Chen et al., 2020), assessments of the climate impacts of aviation (D. Lee et al.,
 210 2010; Schumann et al., 2021), evaluations of radiative transfer in chemistry–climate mod-
 211 els (Forster et al., 2011), and cloud clearing for volcanic plumes (Kylling et al., 2015).

212 Our radiative transfer simulations use the DISORT (discrete ordinate) solver (Stamnes
 213 et al., 1988, 2000), which adopts a one-dimensional geometry under the plane parallel
 214 approximation to calculate radiative transfer across the solar spectrum. Reptran absorp-
 215 tion parameterizations (Gasteiger et al., 2014) are used in the ‘coarse’ configuration, which
 216 corresponds to a spectral range of 240 nm–5 μm and a spectral resolution of 15 cm^{-1} (Emde
 217 et al., 2016), with approximately 5000 bands and 7500 wavelengths represented. Atmo-
 218 spheric background data include height, pressure, temperature, air density, ozone, oxy-

219 gen, water vapor, and CO₂. All data inputs are based on MERRA-2 model-level fields
 220 except for oxygen, which is taken from the libRadtran tropical default profile, and aerosol
 221 profiles from other products as indicated in section 4.2. Unless otherwise specified, so-
 222 lar zenith angle for radiative heating calculations is set to 0° for idealized simulations and
 223 the daily minimum at the corresponding latitude for geolocated simulations, while sur-
 224 face albedo is set to the local mean from MERRA-2 (0.15 for most calculations). These
 225 settings are both simple and helpful for emphasizing ATAL effects on radiative heating,
 226 which, as shown below, are largely insensitive to these two parameters. ATAL effects on
 227 TOA and surface fluxes are sensitive to these parameters, and are therefore computed
 228 in most scenarios by integrating over a representative diurnal cycle for Dhaka, Bangladesh.

229 Aerosol are specified according to OPAC as insoluble (INSO: hydrophobic organic
 230 carbon), water soluble (WASO: sulfate, nitrate, ammonium, hydrophilic black carbon,
 231 and hydrophilic organic carbon), or soot (SOOT: hydrophobic black carbon). To eval-
 232 uate the impacts of including dust in the calculations, the accumulation and coarse modes
 233 of mineral dust (MIAM and MICM) are used according to the size bins simulated by MERRA-
 234 2. The benchmark profile for the control, or ‘base’, simulation is taken from mean con-
 235 ditions for July–September 2011–2020 within 22.5°N–25°N and 87.5°E–90°E, and excludes
 236 dust. Idealized simulations are then conducted to evaluate the impacts of changing the
 237 amplitude, composition, or peak height of the ATAL on clear-sky shortwave surface fluxes,
 238 TOA fluxes, and heating rates around the tropopause. We also evaluate changes in the
 239 solar zenith angle and surface albedo in an idealized setting and over typical diurnal and
 240 seasonal cycles. All idealized simulations, including the base case, only consider aerosol
 241 within the 60–180 hPa layer. This layer definition is based on many previous observa-
 242 tions of ATAL aerosol loading, which place the ATAL between about 13–18 km (360–420 K
 243 potential temperature Vernier et al., 2011, 2017; Brunamonti et al., 2018; Hanumanthu
 244 et al., 2020; Zhang et al., 2020).

245 **3 Aerosol layer description and idealized shortwave effects**

246 **3.1 ATAL distribution and composition in MERRA-2**

247 Figure 1 shows distributions of sulfate, organic carbon, and black carbon within
 248 the UTLS (300 hPa–55 hPa) above the Asian monsoon region. Although dust accounts
 249 for the largest fraction by mass in the MERRA-2 UTLS (see also W. K. M. Lau et al.,
 250 2018), dust concentrations are weighted toward lower altitudes and the northern part
 251 of the anticyclone (Fig. S1 in Supporting information). The distribution of dust during
 252 July–September is consistent with dust transport from source regions in North Africa,
 253 the Middle East, and Central Asia in the subtropical westerly jet being entrained into
 254 the monsoon anticyclone and then lifted isentropically around the eastern flank of the
 255 anticyclone. Comparison with observational estimates (e.g., Vernier et al., 2017) and other
 256 datasets (Fig. S2 in Supporting Information) suggests that MERRA-2 overestimates dust
 257 concentrations at these levels. Moreover, our offline radiative transfer calculations (Fig. 5a)
 258 show that including dust at ATAL altitudes (60–180 hPa) has little impact on clear-sky
 259 shortwave heating rates. We therefore omit it from most of the following analysis.

260 ATAL aerosols are spread throughout the upper-level monsoon anticyclone at 100 hPa
 261 (Fig. 1a), with the largest concentrations along the southern flank near 370 K–375 K po-
 262 tential temperature (~ 110 hPa; Fig. 1a–c). This distribution can be attributed to two
 263 key factors. First, the main convective sources of the ATAL as identified for MERRA-
 264 2 by W. K. M. Lau et al. (2018) are located upstream of the elongated maximum along
 265 the southern flank. Second, strong baroclinicity across the northern part of the anticy-
 266 clone (the subtropical westerly jet) locates potential temperature surfaces at higher pres-
 267 sures (lower altitudes) in the north than in the south, while mean diabatic heating is pos-
 268 itive but weak outside of deep convection (~ 0.5 K day⁻¹; Tegtmeier et al., 2022, their
 269 Figs. 8.59–8.60). Transport within the anticyclone is mostly quasi-isentropic, with ascent

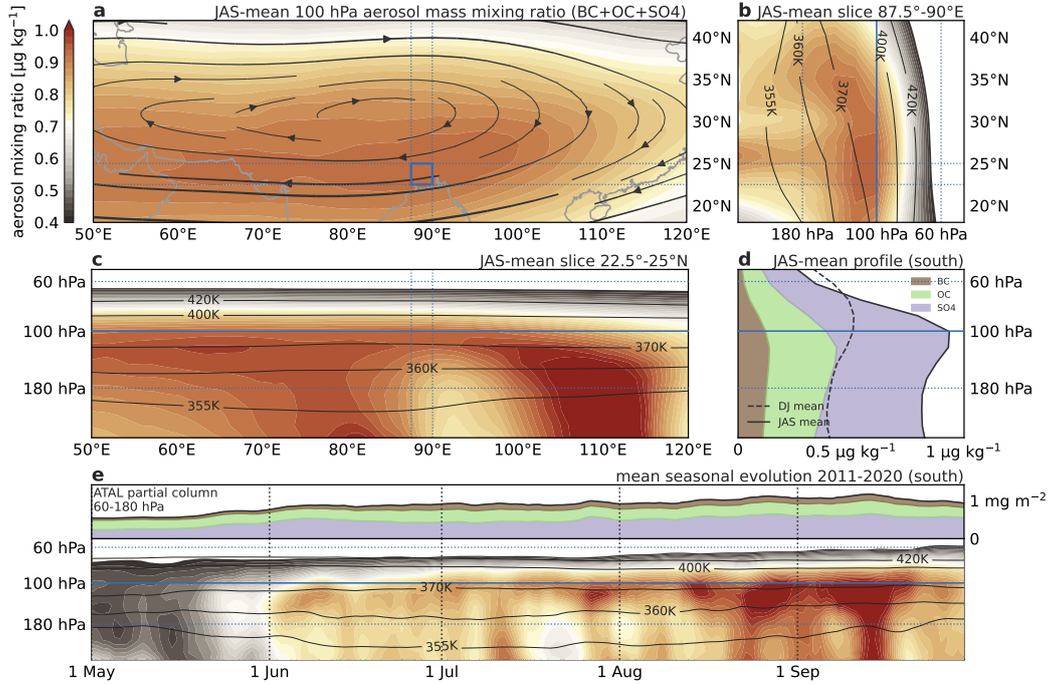


Figure 1. Distributions of sulfate, organic carbon, and black carbon aerosol mass mixing ratios based on the MERRA-2 aerosol reanalysis (a) as a function of latitude and longitude on the 100 hPa isobaric surface, (b) as a function of latitude and pressure along the 87.5°E–90°E longitude band, (c) as a function of longitude and pressure along the 22.5°N–25°N latitude band, and (d) as an area-average profile within 87.5°E–90°E and 22.5°N–25°N for July, August, and September 2011–2020. (e) Mean evolution of aerosol profile (lower panel) and partial column (vertically-integrated over 180–60 hPa; upper panel) within 87.5°E–90°E and 22.5°N–25°N from 1 May to 30 September 2011–2020. Streamlines in (a) show the upper-level anticyclone at 100 hPa based on MERRA-2. Contours in (b), (c), and (e) show potential temperature surfaces spanning the upper troposphere and lower stratosphere. Dashed line in (d) represents sum of sulfate, black carbon and organic carbon in December 2007 – January 2008. Shaded regions in (d) and (e) illustrate the relative abundances of black carbon (brown), organic carbon (green), and sulfate (purple) aerosol.

270 upward along isentropes in the east (where flow is north-to-south) and descent in the west
 271 (where flow is south-to-north). The isobaric aerosol distribution therefore shows larger
 272 values where the 370 K–375 K isentropic layer outcrops.

273 Cross-sections of mean ATAL vertical structure over July–September 2011–2020
 274 are shown for the north-south (averaged zonally across 87.5°E–90°E; Fig. 1b) and east-
 275 west (averaged meridionally across 22.5°N–25°N) directions. The north-south cross-section
 276 cuts across the nominal center of the anticyclone above the eastern Tibetan Plateau, while
 277 the east-west cross-section slices eastward along the southern flank of the anticyclone (Fig. 1a).
 278 These cross-sections show a clearly defined ATAL peaking near the 370 K potential tem-
 279 perature surface (~ 110 hPa along the east-west cross-section). This enhanced aerosol layer
 280 is fed by convective uplift between 20°N and 30°N with centers within 70°E–90°E (North
 281 India) and 105°E–115°E (Sichuan Basin and southern China; Fig. 1c). The aerosol max-
 282 imum in the north-south cross-section largely follows isentropic contours (Fig. 1b) de-
 283 spite long transport distances around the anticyclone from the convective source regions.
 284 Enhanced aerosol loading in this altitude range develops mainly from the beginning of
 285 June and persists through the end of September, with the largest values in late August
 286 and early September.

287 Compared with W. K. M. Lau et al. (2018), our results suggest an additional ‘chim-
 288 ney’ over southern China ($\sim 110^\circ\text{E}$; Fig. 1c), which is broader and shifted to the south-
 289 east relative to the Sichuan Basin source highlighted in their results. This peak could
 290 indicate persistent southeastward transport from the Sichuan Basin; however, the mean
 291 upper-level flow from the Sichuan Basin is southwestward (see also K.-O. Lee et al., 2021).
 292 We note further that W. K. M. Lau et al. (2018) analyzed only one monsoon season (May–
 293 August 2008) and did not include sulfate. There are thus several possible reasons for the
 294 difference between our results and theirs. First, it could be due to interannual variabil-
 295 ity in convective sources over East Asia (e.g., strong influences of convection over south-
 296 ern China during summer 2017; Bucci et al., 2020). Second, although local maxima in
 297 OC and BC are evident in that region (Figs. S3 and S4 in Supporting Information), they
 298 are much weaker than those over South Asia, and the chimney-like connection to the tropo-
 299 sphere as in Fig. 1 appears only for sulfate (Fig. S5 in Supporting Information). Fi-
 300 nally, the mean seasonal cycle in our core region, downstream of convection over south-
 301 ern China, suggests that sulfate loading in the ATAL is relatively small through most
 302 of July before increasing in August and peaking in early September (Fig. S5e in Sup-
 303 porting Information). This seasonality may reflect a strengthening convective source over
 304 southern China as the East Asian monsoon rainband retreats, changes in the efficiency
 305 of sulfate removal by deep convection over the Bay of Bengal (which starts early and peaks
 306 in July), or some combination of the two.

307 Within the MERRA-2 ATAL as defined in this work (60–180 hPa), sulfate is the
 308 largest component by mass fraction (43–61% for the ATAL column over the seasonal cy-
 309 cle; upper part of Fig. 1e). OC is next largest (28%–48%), with BC mass fractions smaller
 310 than OC by about a factor three (9%–16%). Notably, the hydrophilic component of OC
 311 is approximately in steady state through the monsoon development cycle (Fig. S4e), with
 312 the OC column almost completely hydrophilic in May and subsequent changes dominated
 313 by changes in the hydrophobic component. Here the ATAL column is obtained by in-
 314 tegrating across the depth of the layer in pressure coordinates:

$$C_{\text{ATAL}} = \frac{1}{g} \int_{p_b}^{p_t} q_{\text{aer}} dp \quad (1)$$

315 where $g = 9.8 \text{ m s}^{-2}$ is the gravitational acceleration, $p_b = 180 \text{ hPa}$ is the base of the
 316 layer, $p_t = 60 \text{ hPa}$ is the top of the layer, and q_{aer} is the aerosol mass mixing ratio. The
 317 OC and BC column masses calculated in this way are highly correlated ($r = 0.75$) but
 318 the majority of this is in the spatial dimension ($r = 0.88$), with much smaller covari-
 319 ability in time ($r = 0.59$). OC and sulfate are also highly correlated ($r = 0.64$), with

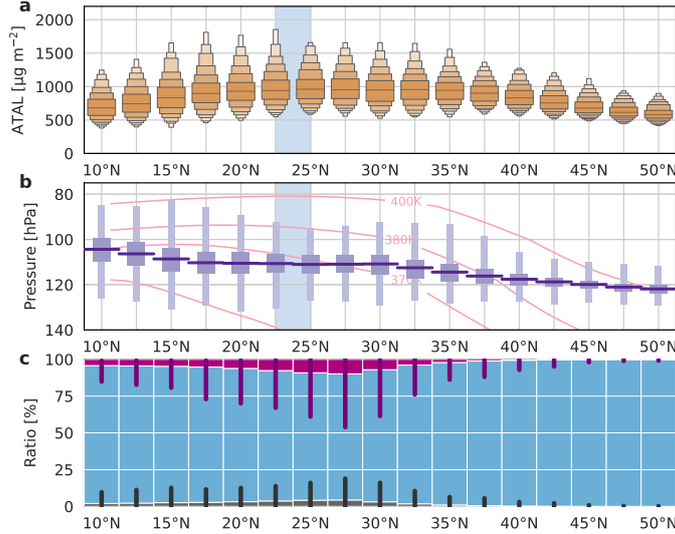


Figure 2. Variations in latitude of (a) distributions of daily-mean vertically-integrated aerosol mass per unit area in the 60–180 hPa layer; (b) box-and-whisker plots of ATAL height (in pressure) defined as the daily-mean center-of-mass (50% of ATAL mass both above and below); and (c) the soot (grey), water soluble (blue), and insoluble (purple) fractions of ATAL aerosol in 2.5° latitude bins along the 87.5°E–90°E longitude band. Minimum and maximum soot and insoluble fractions are indicated by thin grey and purple bars, respectively. Pink contours in (b) show potential temperature surfaces.

roughly equal correlations in space and time, while BC and sulfate are only weakly correlated ($r = 0.36$), especially in time ($r = 0.22$). Similar ranges of relative concentration are obtained for the mean vertical profile over July–September (Fig. 1d), with sulfate ratios ranging from 50% near the base of the layer at 180 hPa to 60% near the top of the layer at 60 hPa. OC mass fractions are consistently around 30% (28–33%) through the depth of the ATAL, while BC mass fractions decrease by half from the base of the layer (16%) to the top of the layer (8%). Given the weak temporal correlations and different vertical distributions of BC and sulfate within the ATAL, we treat these two species as nominally independent in setting up the idealized simulations below.

Figure 2 displays summary information about variations in selected ATAL properties by latitude along the same north-south cross-section as Fig. 1b. ATAL amplitude (Fig. 2a), calculated daily according to equation 1, peaks along the southern flank of the anticyclone around 20°N–25°N (Fig. 2a). The largest variations in daily-mean amplitude are located slightly south of the peak values (15°N–22.5°N) where, despite smaller median values, large outliers on the high end indicate episodes of strong convective uplift of polluted air. Variance reduces sharply with latitude northward of 35°N.

We estimate the daily height of the aerosol layer as its ‘center of mass’. This is calculated by first integrating aerosol partial columns:

$$C(p) = \frac{1}{g} \int_p^{p_t} q_{\text{aer}} dp \quad (2)$$

for $p_b \leq p < p_t$, then calculating the ratio of $C(p)$ relative to C_{ATAL} (equation 1), and finally linearly interpolating the ratio in $\ln(p)$ to find the pressure associated with the ratio 0.5 (p_{com}). The results are not qualitatively sensitive to ratios between 0.25 and 0.75. Defined in this way, ATAL height decreases (p_{com} increases) from south to north,

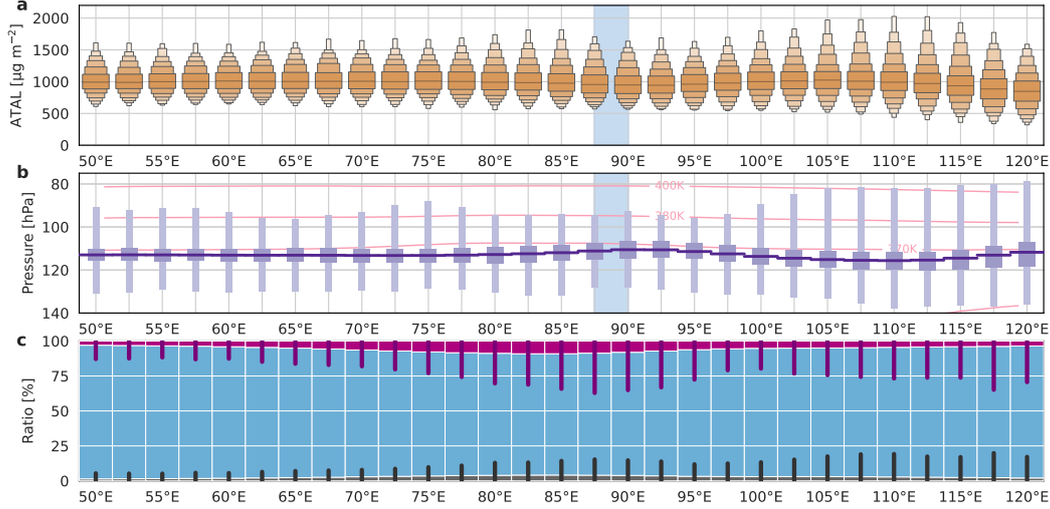


Figure 3. As in Fig. 2 but for variations in longitude within the 22.5°N–25°N latitude band.

342 with the largest variability at low latitudes. Daily-mean heights frequently lie above 370 K
 343 at all latitudes, with a few examples even reaching 400 K. ATAL heights are very con-
 344 sistent across the southern part of the anticyclone (20°N–30°N). Lower heights north of
 345 30°N can be explained at least in part by the northward decline of isentropic surfaces
 346 toward higher pressures.

347 Relative composition of the mean ATAL layer is shown in Fig. 2c, here converted
 348 from the MERRA-2 species classification (sulfate, OC, BC) to the OPAC optical prop-
 349 erties classification (WASO, INSO, SOOT). WASO is the largest fraction through the
 350 entire domain, consisting of sulfate and the hydrophilic components of OC and BC. SOOT
 351 (hydrophobic BC) and INSO (hydrophobic OC) fractions increase northward from low
 352 latitudes to 27.5°N, above the south slope of the Himalayas, and then decrease north-
 353 ward to the northern flank of the anticyclone around 40°N (the subtropical westerly jet).
 354 Distributions north of 40°N show little variability and are almost completely composed
 355 of sulfate. We therefore assume these distributions to be representative of extratropi-
 356 cal stratospheric background aerosol and omit them from further analysis.

357 Average ATAL amplitude is fairly consistent as a function of longitude within the
 358 22.5°N–25°N latitude band (Fig. 3a), although peaks in variance are evident around the
 359 two convective centers over North India (80°E–85°E) and southern China (105°E–115°E).
 360 Variance is especially large over the latter region, where the largest values of daily-mean
 361 C_{ATAL} exceed $2000 \mu\text{g m}^{-2}$. Variations in ATAL height across longitudes (Fig. 3b) ap-
 362 pear to be linked mainly to dilution and enhanced removal of the lower part of the layer
 363 by strong convection over the Bay of Bengal (near 90°E) and South China Sea (near 120°E).
 364 Composition fractions are consistent with the hypothesis that convection over North In-
 365 dia (and the Sichuan Basin upstream) represent the main sources of carbonaceous aerosols,
 366 as the INSO and SOOT fractions are largest westward of 100°E. By contrast, the con-
 367 vective source over southern China has little impact on the relative concentrations of INSO
 368 and SOOT, consistent with this source providing mainly water soluble aerosols to the
 369 ATAL as represented in MERRA-2.

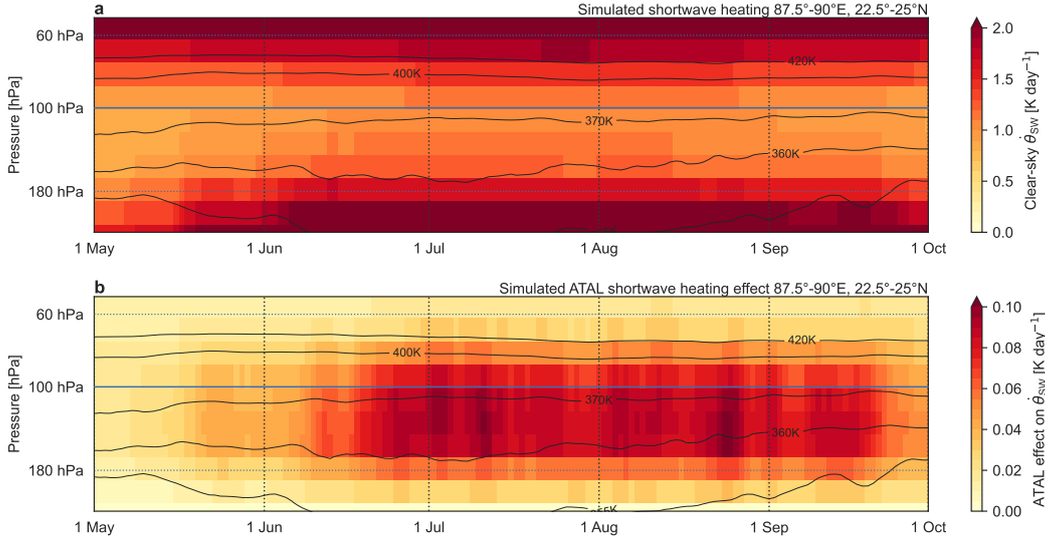


Figure 4. Mean seasonal evolution of (a) clear-sky shortwave heating and (b) the ATAL effect on clear-sky shortwave heating based on the MERRA-2 aerosol reanalysis. Radiative heating is calculated using offline radiative transfer simulations that account for seasonal changes in temperature, water vapor, ozone, ATAL aerosol (within 60–180 hPa), surface albedo, and the daily minimum solar zenith angle during 1 May–30 September 2011–2020. Heating rates have been divided by the Exner function to convert $\partial T/\partial t$ to $\dot{\theta}$ for ease of comparison to potential temperature contours.

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3.2 Idealized simulations of ATAL effects on radiative heating

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In this section, we summarize the results of offline radiative transfer calculations targeting different properties of the ATAL (Table 1). For context, Figure 4 shows the mean seasonal cycle of simulated shortwave heating for a clear-sky atmosphere with aerosol above the core region (22.5°N–25°N, 87.5°N–90°N), along with the aerosol contribution to this heating based on the evolution of ATAL aerosols shown in Fig. 1e. The ATAL radiative effect on clear-sky shortwave heating in this region grows slowly from mid-May before increasing sharply in mid-June (Fig. 4b). Aerosol effects on clear-sky shortwave heating persist at comparable levels until late September. These effects are strongest between 360 K and 400 K potential temperature, where clear-sky shortwave heating is at a minimum (Fig. 4a). Overall, the magnitude of the aerosol effect adds an additional heating of roughly 10% to clear-sky shortwave heating in the 100–150 hPa layer during the peak monsoon season. Evaluating ATAL effects relative to a wintertime aerosol profile rather than the no-aerosol profile reduces radiative heating effects by about half (Fig. S6).

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Figure 5 shows aerosol effects on the surface flux, TOA flux, and radiative heating in the UTLS from the perspectives of ATAL amplitude, surface albedo, and solar zenith angle. The offline calculation for the base profile (i.e., adopting mean conditions for the core region) is compared to an offline calculation without aerosol and the online calculation from MERRA-2 in Figure 5a. The general structure of the profile is consistent with that from MERRA-2, but values are slightly smaller through most of the UTLS. These differences could result from nonlinearities in how the effects of shortwave absorbers scale; i.e., comparing an offline radiative calculation based on mean conditions (for which the cold point, hygropause, and ozone gradient are fixed in the vertical) to a mean of online profiles calculated from instantaneous conditions (for which the cold point, hygropause,

Table 1. Summary description of idealized radiative transfer calculations. All simulations use the same July–September (JAS) 2011–2020 mean atmospheric profile for 22.5°N–25°N and 87.5°E–90°E from MERRA-2. The base aerosol profile is similarly defined as JAS-mean BC+OC+SO₄ within 60–180 hPa. C_{ATAL} refers to aerosol column mass (eq. 1); Prf_{ATAL} to the variation of aerosol type mass fractions with height; WASO, INSO, and SOOT to mass fractions of each aerosol type; SZA to solar zenith angle; and ALB to surface albedo.

Experiment	C_{ATAL}	Prf_{ATAL}	WASO	INSO	SOOT	SZA ^a	ALB ^a	Figure
amplitude	$n \times \text{base}^b$	base	base	base	base	0°	0.15	Fig. 5b
solar	base	base	base	base	base	0–80°	0–0.25	Fig. 5c
diurnal	$n \times \text{base}^b$	base	base	base	base	15 Aug ^a	15 Aug ^a	Fig. 6
composition	base	constant	0.6–1	0–0.4	0–0.2	0°	0.15	Fig. 7
height	selected ^c	base	selected	selected	selected	0°	0.15	Fig. 8
validation	product	product	product	product	product	0°	0.15	Fig. 12

^a Solar parameters for TOA and surface fluxes specified as for Dhaka, Bangladesh on 15 August.

^b n is a multiplier between 0.1 and 4.

^c Daily mean profiles in each height bin are randomly selected from July–September 2011–2020.

394 and ozone gradient may vary in the vertical). They may also result from different assump-
 395 tions about aerosol optical properties, as our conservative approach (assigning hydropho-
 396 bic OC to INSO and hydrophilic OC and BC to WASO) minimizes the absorbing aerosol
 397 fraction. As ATAL effects scale linearly with amplitude and the relative composition of
 398 absorbing versus scattering aerosols (see below), we take the simulated aerosol effect from
 399 libRadtran as representative while accounting for these possible biases when computing
 400 relative effects.

401 Increasing the solar zenith angle from 0° to 40° has little impact on the ATAL ef-
 402 fect on shortwave heating (Fig. 5c) owing to compensating effects between decreases in
 403 TOA insolation (proportional to the cosine of solar zenith angle) and increases in path
 404 length (inversely proportional to the cosine) as solar zenith angle increases. The heat-
 405 ing effect weakens substantially as the solar zenith angle approaches 90°. Increasing sur-
 406 face albedo increases the ATAL effect on clear-sky shortwave heating, but these differ-
 407 ences are small and require relatively large changes in the albedo. This sensitivity can
 408 be safely ignored for the clear-sky case in our core region as variations in surface albedo
 409 are small, but may be influential when the aerosol layer overlies thick anvil clouds or land
 410 ice, which are both present in abundance within the Asian monsoon domain. Changes
 411 in the no-aerosol radiative heating rates associated with changes in albedo and solar zenith
 412 angle are not shown, but are accounted for when computing the ATAL radiative effects.

413 Further context is provided by evaluating ATAL radiative effects across a repre-
 414 sentative diurnal cycle (Fig. 6). Integrated over the day, the upward flux at the nom-
 415 inal top-of-atmosphere (TOA) increases by about 0.04 W m^{-2} , while the net downward
 416 (absorbed) flux of solar radiation at the surface decreases by about 0.32 W m^{-2} (Fig. 6a-
 417 b). The TOA radiative effect is about one third of that reported by Vernier et al. (2015),
 418 who estimated this effect to 0.12 W m^{-2} . The difference between our estimate and theirs
 419 is further enhanced if we follow their approach and adopt a representative winter aerosol
 420 profile (dashed line in Fig. 1d) as the baseline in place of the no-aerosol case. With
 421 the same diurnal cycle of solar parameters, the decrease in surface absorption is reduced to
 422 0.19 W m^{-2} while the change in the upward flux at TOA reverses sign to -0.04 W m^{-2} .
 423 The larger decrease in the surface effect relative to TOA implies a reduction in the ATAL
 424 heating, as seen also in Fig. S6 in comparison to Fig. 4. The difference in TOA effect in
 425 our calculations relative to that reported by Vernier et al. (2015) can be largely attributed

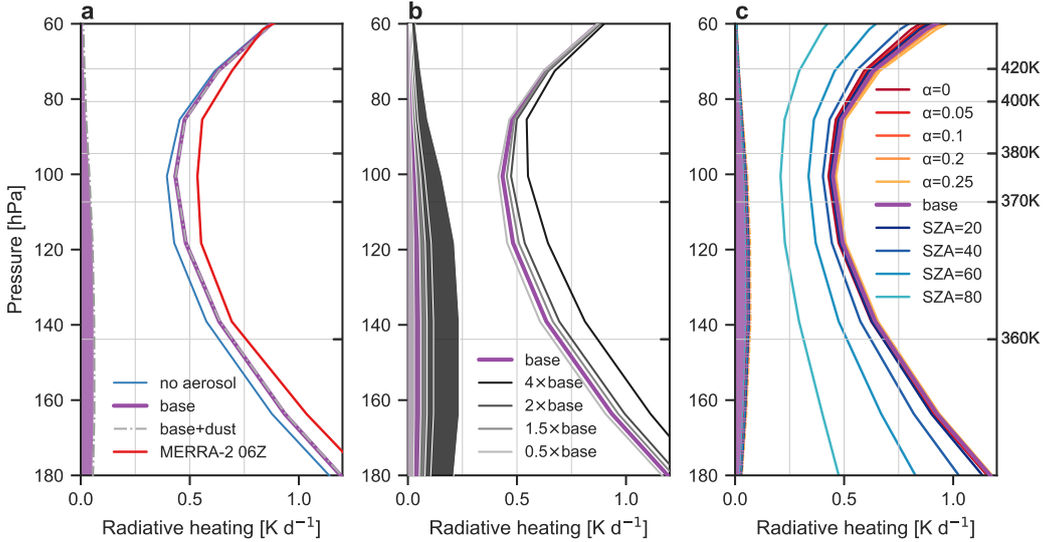


Figure 5. Radiative heating profiles calculated for (a) the no-aerosol (blue), base (purple; ATAL aerosol as in Fig. 1d within 60–180 hPa only), and base+dust (grey dash-dot line) profiles; (b) different ATAL amplitudes ranging from 0.5×base to 4×base without changing relative composition; and (c) for the base profile at selected values of solar zenith angle (SZA) and albedo (α); profiles in panels a and b are based on SZA = 0 and $\alpha = 0.15$. MERRA-2 clear-sky shortwave heating for local mid-day (04:30–07:30 UTC) is shown in c for comparison. Input data other than aerosols are specified as mean values for JAS 2011–2020 in 22.5°N–25°N and 87.5°E–90°E.

426 to the inclusion of BC (only OC and sulfate were considered in the calculations by Vernier
 427 et al., 2015), which enhances atmospheric absorption. We can retrieve a TOA effect of
 428 0.12 W m^{-2} by raising the WASO fraction to 96% or higher (Fig. 7a), lowering the SOOT
 429 fraction to less than 2% (Fig. 7c), or increasing the base ATAL amplitude by a factor
 430 3 (Fig. 6a). This last result underscores the linearity of the ATAL effect at both TOA
 431 and surface. Moreover, it highlights the relatively weak radiative impacts of dust within
 432 the ATAL. Including dust within the ATAL as represented by MERRA-2 for the core
 433 region increases the amplitude (i.e., the column mass per unit area) by a factor of 2.8,
 434 but only alters the TOA and surface effects by $\sim 10\%$, far less than scaling the base
 435 amplitude by a similar amount. As the additional decrease in surface absorption due to dust
 436 ($\sim 0.03 \text{ W m}^{-2}$) is partially compensated by increased upward flux at TOA ($\sim 0.01 \text{ W m}^{-2}$),
 437 its effect on radiative heating within the ATAL vertical range is very weak (Fig. 5a).

438 Diurnal variations in the ATAL effect on clear-sky shortwave heating are small over
 439 daylight hours (Fig. 6d). The largest sensitivities are found for the middle of the day,
 440 but effects are comparable in magnitude during the daytime except just after sunrise and
 441 just before sunset. Accordingly, given ~ 13 hours of daylight, the mean ATAL effect on
 442 clear-sky shortwave heating integrated over 24 h is approximately half of that calculated
 443 for mid-day. To better emphasize the ATAL forcing on shortwave heating during day-
 444 light hours, we adopt a solar zenith angle of 0° for all further calculations and compar-
 445 isons of radiative heating. By contrast, flux effects at TOA and surface are all integrated
 446 over 24 h to facilitate comparison with previous work.

447 We evaluate the effects of ATAL composition by holding the total concentration
 448 of anthropogenic aerosols fixed while changing the proportions of different species (Fig. 7).
 449 The ratios of all three species are vertically homogeneous in each experiment included

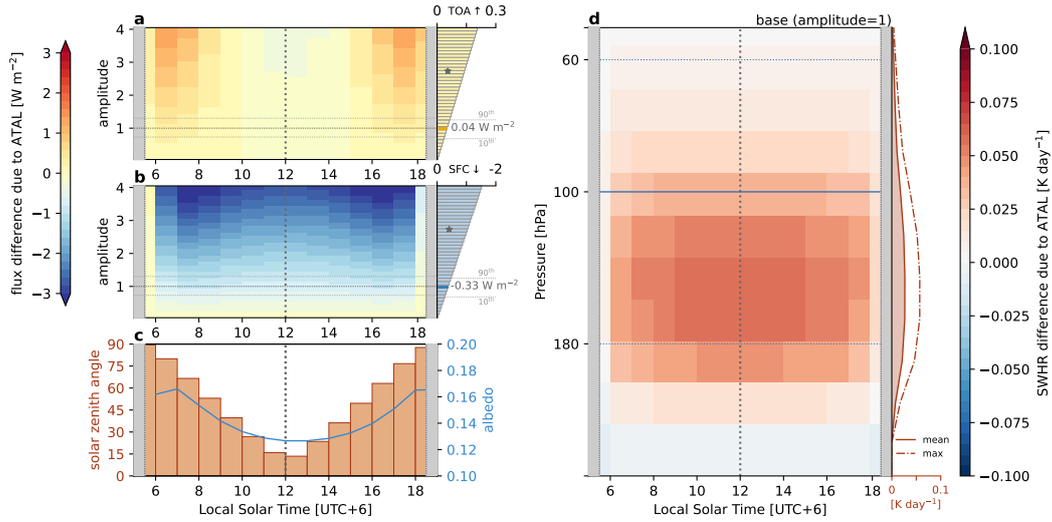


Figure 6. Diurnal variations of ATAL effects on clear-sky shortwave fluxes at (a) the top-of-atmosphere (TOA) and (b) clear-sky shortwave fluxes at the surface for ATAL amplitudes ranging from $0.1 \times \text{base}$ to $4 \times \text{base}$; (c) diurnal variations in albedo (line) and solar zenith angle (bars), and (d) diurnal variations in ATAL effects on clear-sky shortwave heating for the ATAL base profile (aerosol within 60–180 hPa distributed as in Fig. 1d). All effects are calculated relative to no-aerosol simulations with matching solar parameters. Daily-mean values for ATAL effects on TOA flux, surface flux, and radiative heating are shown along the right margins of the corresponding panels, with values for the base profile highlighted. The 10th and 90th percentiles of ATAL column mass (eq. 1) for the core region in MERRA-2 are marked, as are the TOA and surface effects of including dust in addition to the base ATAL (grey stars).

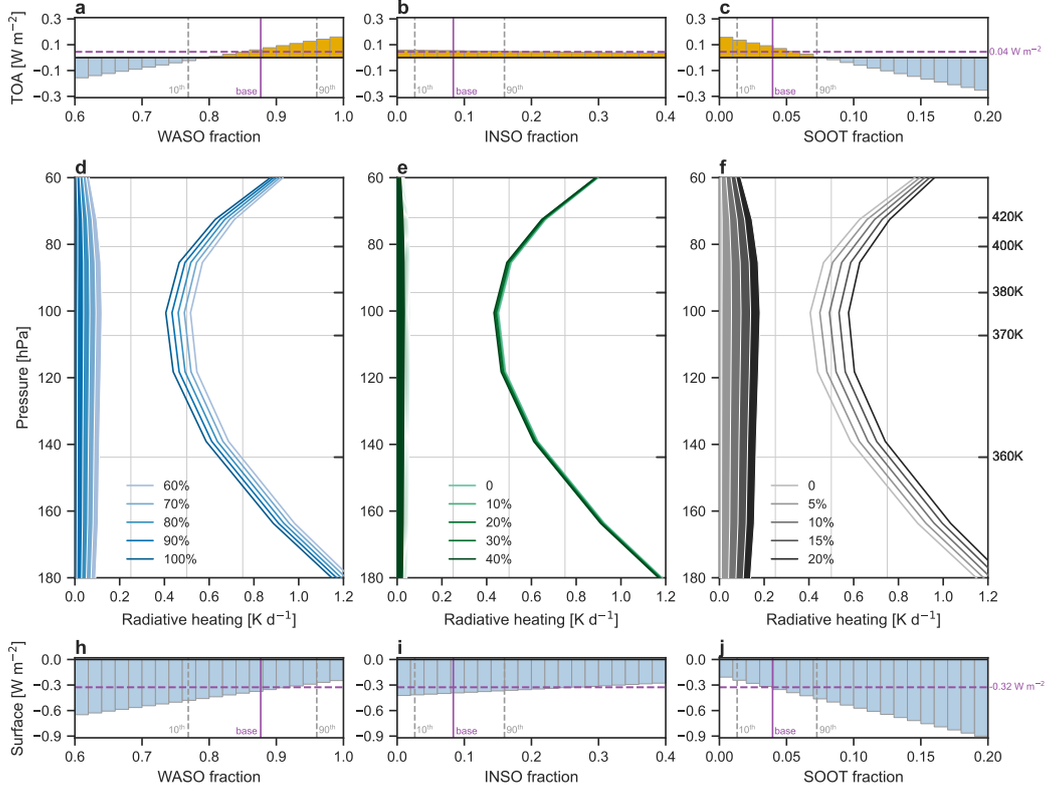


Figure 7. Variations in (a)–(c) top-of-atmosphere (TOA) shortwave flux anomalies, (d)–(f) vertical profiles of shortwave radiative heating in the tropopause layer, and (h)–(j) net surface shortwave flux anomalies under clear-sky conditions as a function of variations in the fractions of (a,d,h) water-soluble, (b,e,i) insoluble, and (c,f,j) soot relative to total aerosol. Mass fractions of each aerosol type are constant in height for each experiment. Heating rate profiles are calculated assuming a solar zenith angle of zero and a surface albedo of 0.15; TOA and surface fluxes are daily-mean values as in Fig. 6. Differences in heating relative to the no-aerosol case are shown as shaded regions on the left of panels d–f. The 10th and 90th percentiles of daily-mean ATAL column mass in the core region are marked as grey dashed vertical lines in panels 1–c and h–j, along with the base column mass (vertical purple line) and the TOA (a–c) or surface (h–j) flux effect calculated for base (dashed horizontal purple line).

450 in Fig. 7, resulting in different vertical structures relative to the base profile. Among the
 451 three OPAC aerosol types (WASO, INSO, SOOT), SOOT has the largest impact on ra-
 452 diative heating despite its small proportion (Fig. 7c). This large effect on clear-sky short-
 453 wave heating is consistent with its large capacity to absorb solar radiation (Samset et
 454 al., 2018). Compared to the other two types, SOOT has larger extinction coefficients across
 455 the solar spectrum and a consistently small single-scattering albedo, averaging ~ 0.2 for
 456 wavelengths less than $1 \mu\text{m}$ (Hess et al., 1998). Extinction coefficients for WASO are ap-
 457 proximately half of those for SOOT but with single-scattering albedos close to 1, while
 458 extinction coefficients for INSO are small by comparison ($\sim 10\%$ of those for WASO) with
 459 single-scattering albedos of approximately 0.8. The sensitivities in Fig. 7 may thus be
 460 understood as resulting largely from competition between a large fraction of scattering
 461 aerosols with moderate extinction coefficients and a small fraction of absorbing aerosols
 462 with large extinction coefficients. Increases in the former result in larger backscattering
 463 to the TOA (Fig. 7a), reduced ATAL heating near the tropopause (Fig. 7d), and a weaker
 464 reduction of net downward flux at the surface (Fig. 7h). Increases in the latter result in
 465 smaller backscattering (Fig. 7c), enhanced ATAL heating near the tropopause (Fig. 7f),
 466 and a stronger reduction of net downward flux at the surface (Fig. 7j). Although the INSO
 467 component is about twice as large as that for SOOT, small extinction coefficients mean
 468 that its impact in these scenarios is mainly to proportionately reduce both the absorb-
 469 ing and scattering effects while having little radiative impact itself. Note that brown car-
 470 bon, as the absorbing component of OC (Samset et al., 2018), is not considered in these
 471 simulations, and hydrophilic BC is apportioned to WASO. Increasing the SOOT frac-
 472 tion as in Fig. 7f may thus be considered as a crude approximation to partitioning part
 473 of the hydrophobic OC to SOOT rather than INSO, or as retaining the hydrophilic frac-
 474 tion of BC in SOOT (Rémy et al., 2019).

475 Figure 8 shows results from sensitivity experiments evaluating the effects of vary-
 476 ing the ATAL height. Here, height is defined as the ‘center-of-mass’ pressure level (p_{com})
 477 for which 50% of the ATAL column by mass is located both between 60 hPa and p_{com}
 478 and between p_{com} and 180 hPa. For reference, p_{com} for the base profile is 112 hPa (Fig. 8c).
 479 Six height bins are defined, each with ten randomly selected daily-mean profiles for which
 480 p_{com} is within ± 0.5 hPa of the specified level. Although total aerosol mass is the same
 481 in all simulations (Table 1), ATAL vertical structure (Fig. 8c) and composition (Fig. 8a-
 482 b) show substantial differences even within individual bins. These differences illustrate
 483 the scale of day-to-day variability in the ATAL as represented by MERRA-2. No clear
 484 dependence on ATAL height is evident in effects on either the TOA and surface fluxes
 485 or the tropopause-layer radiative heating. Instead, the results highlight the crucial in-
 486 fluence of aerosol composition. Height bins for which a greater number of profiles have
 487 large SOOT mass fractions and small WASO mass fractions show a weaker enhancement
 488 of upward flux at the TOA, a stronger reduction in downward flux at the surface, and
 489 a larger influence on radiative heating near the tropopause. These relationships are repli-
 490 cated within each individual height bin (Fig. 8a-b).

491 The results shown in Fig. 8 exclude interdependence of height and amplitude by
 492 design. These two variables are significantly correlated in this region ($r = 0.62$) so that,
 493 on average, lower heights (larger p_{com}) are associated with a larger aerosol mass per unit
 494 area. We have previously shown that higher ATAL heights are mainly found where mar-
 495 itime deep convection may dilute or wash out the lower part of the aerosol layer (Fig. 1c;
 496 Fig. 3b). The relationships shown in Fig. 5b thus imply that aerosol effects on shortwave
 497 fluxes and heating rates are likely overestimated for height bins with $p_{\text{com}} < 112$ hPa
 498 and underestimated for height bins with $p_{\text{com}} > 112$ hPa in Fig. 8. Simulations based
 499 on varying the height of the peak aerosol concentration with composition fixed as in the
 500 base profile show a weak dependence on height (Fig. S7 in Supporting Information), with
 501 lower peaks associated with a weaker enhancement in upward flux at the TOA, a stronger
 502 reduction of downward shortwave flux, and an increase and downward shift of the max-
 503 imum response in shortwave heating. However, these results also contain the influence

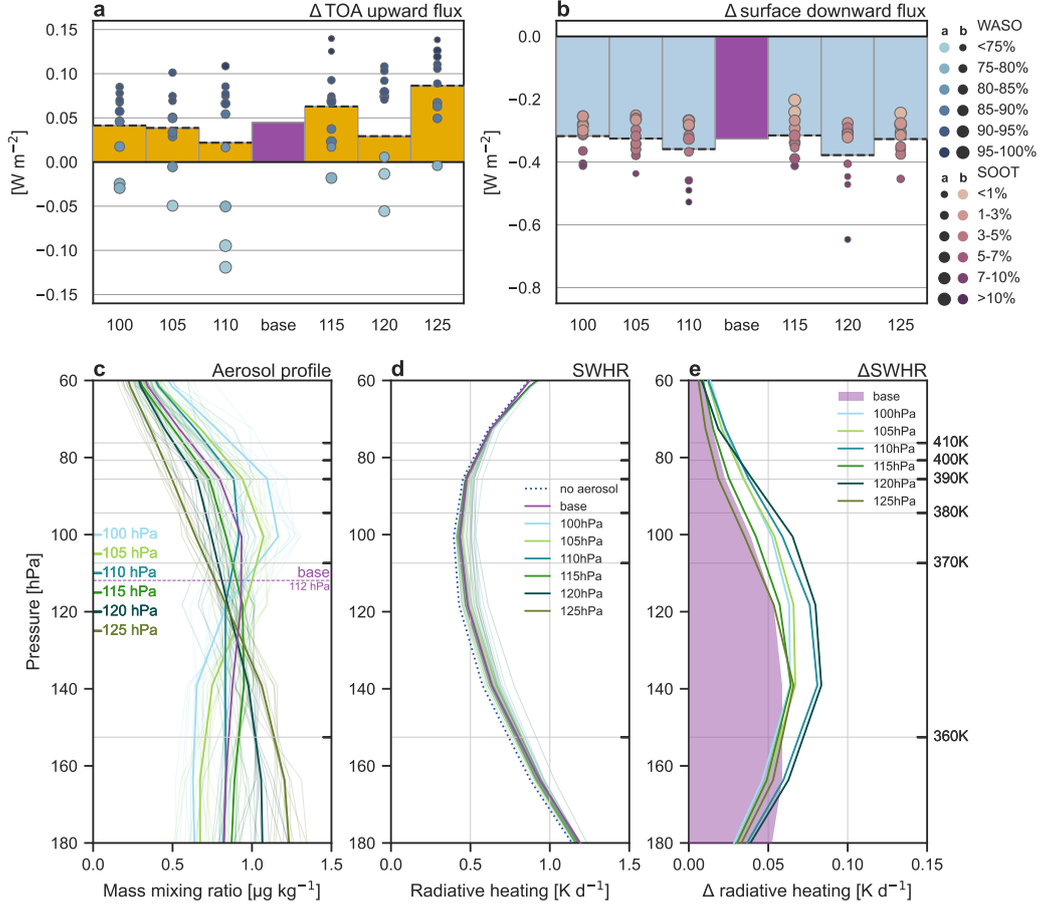


Figure 8. Variations in (a) top-of-atmosphere (TOA) upward shortwave flux anomalies, (b) surface downward shortwave net flux anomalies, (c) vertical profiles of aerosol mass mixing ratio, (d) shortwave radiative heating, and (e) aerosol effects on shortwave radiative heating relative to the no-aerosol case for profiles with different ATAL heights. Data for the 10 randomly selected daily-mean profiles from the core region ($22.5^{\circ}N-25^{\circ}N$, $87.5^{\circ}E-90^{\circ}E$) are shown as round symbols in a-b (key at upper right) and light lines in c-d. Aerosol profiles are normalized to have the same column mass as the base profile in all experiments. Dashed black lines in a and b are included to confirm that mean results based on individual profiles agree well with results based on the mean profile. Heating rate profiles assume a solar zenith angle of zero and a surface albedo of 0.15; TOA and surface fluxes are daily-mean values as in Fig. 6.

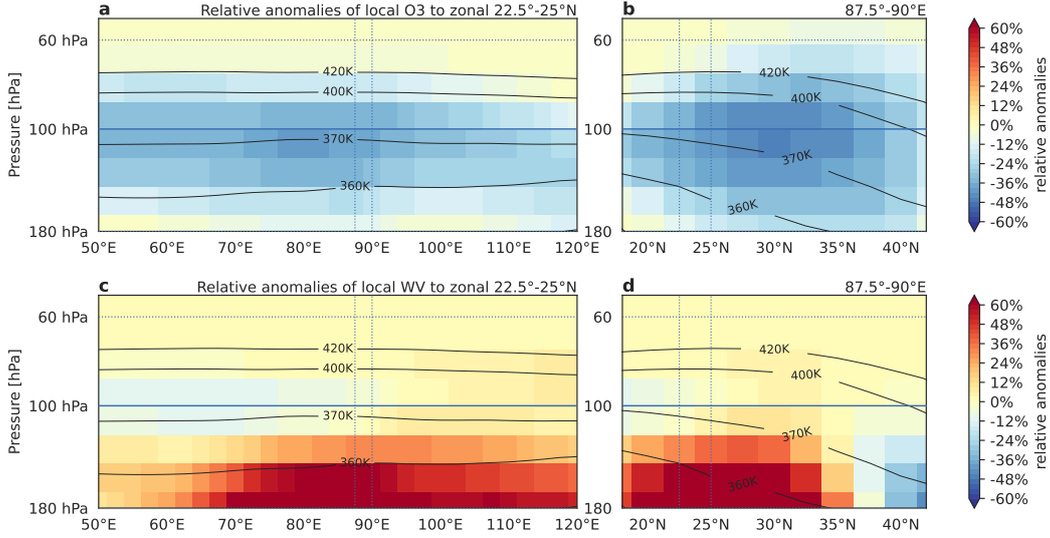


Figure 9. Anomalies in (a)–(b) ozone and (c)–(d) water vapor relative to the corresponding zonal mean profiles based on the MERRA-2 reanalysis for July–September 2011–2020. East–west cross-sections in (a) and (c) are averaged within the 22.5°N–25°N latitude band, while north–south cross-sections in (b) and (d) are averaged within the 87.5°E–90°E longitude band.

504 of changes in composition, as SOOT ratios are largest near the base of the layer while
 505 the top of the layer is almost entirely WASO.

506 4 Aerosol layer shortwave heating effects in context

507 4.1 Comparison to water vapor and ozone effects

508 To place the effects of the ATAL on radiative heating into context, we compare them
 509 to those of monsoon-driven perturbations in water vapor and ozone in the UTLS. When
 510 compared to the zonal mean, upper tropospheric humidity is substantially enhanced above
 511 the monsoon while ozone concentrations are relatively low (e.g., Santee et al., 2017), with
 512 the latter often referred to as an ‘ozone valley’ (e.g., Bian et al., 2020). Figure 9 shows
 513 anomalies in both ozone and water vapor relative to the zonal mean from MERRA-2 re-
 514 analysis products for July–September 2011–2020, again using the zonal and meridional
 515 cross-sections introduced in Fig. 1. Ozone concentrations are about 30% smaller than
 516 the zonal mean within the layer bounded by the 360 K and 420 K potential temperature
 517 surfaces. By contrast, water vapor is enhanced by approximately 30% relative to the zonal
 518 mean in the layer below ~ 360 K, with a weak east–west dipole above. MERRA-2 relaxes
 519 model-generated water vapor concentrations to a zonal-mean climatology above 60 hPa,
 520 so that anomalies are constrained to remain close to zero in the lower stratosphere.

521 Figure 10 shows ATAL effects on radiative heating (Fig. 10a-b) in comparison to
 522 those of ozone (Fig. 10c-d) and water vapor (Fig. 10e-f). Aerosol effects are calculated
 523 relative to the no-aerosol case in each column, as in section 3.2 for the core region. Wa-
 524 ter vapor and ozone effects are calculated by replacing local mean mixing ratios of each
 525 component with zonal mean values at all levels between 60 hPa and 180 hPa. Radiative
 526 transfer calculations are then conducted for the original and perturbed profiles, with ra-
 527 diative effects defined as the difference (original minus perturbed).

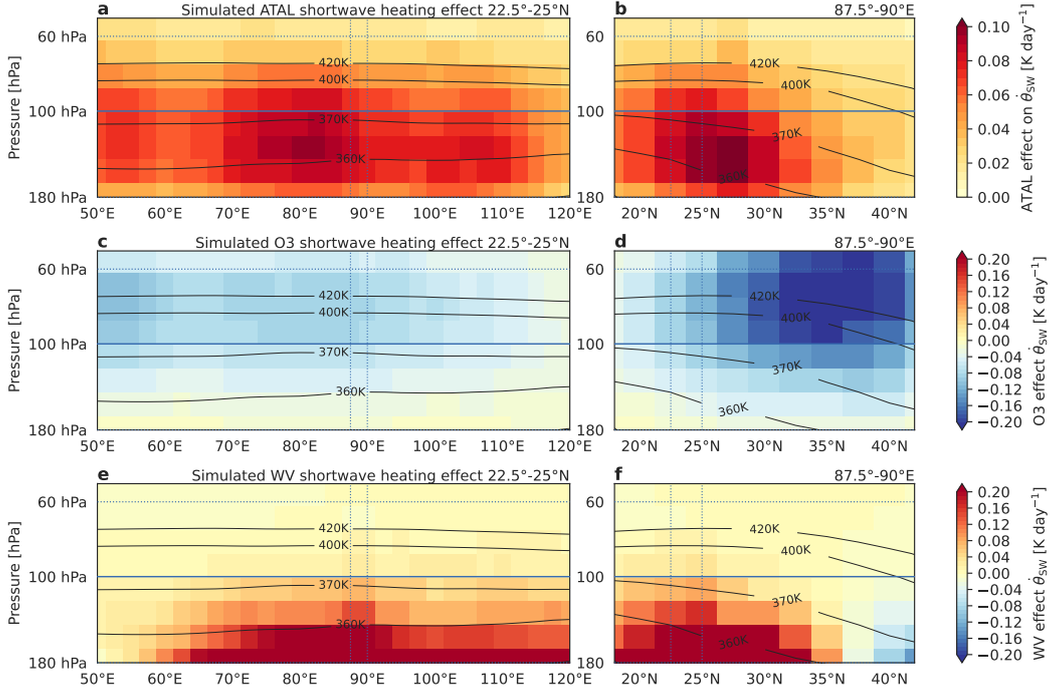


Figure 10. Effects on shortwave radiative heating attributed to (a)–(b) ATAL aerosol (60–180 hPa) relative to the no-aerosol simulation, (c)–(d) ozone anomalies relative to the zonal mean (see Fig. 9a–b) and (e)–(f) water vapor anomalies relative to the zonal mean (see Fig. 9c–d). Radiative effects in panels (a)–(f) are based on offline calculations under clear-sky conditions with surface albedo taken from MERRA-2 and solar zenith angle taken as the zonal minimum on 1 August. Heating rates have been divided by the Exner function to convert $\partial T/\partial t$ to $\dot{\theta}$ for ease of comparison to potential temperature contours.

528 Although maximum values of ozone and water vapor effects within the monsoon
 529 UTLS exceed that of the ATAL effect by about a factor two, values of the ATAL effect
 530 within the 360 K–420 K layer are comparable and often larger in magnitude, especially
 531 along the southern flank of the anticyclone (Fig. 10a–b). The ATAL effect is located at
 532 lower altitudes and is opposite in sign relative to the ozone effect (Fig. 10c–d). Water
 533 vapor effects are concentrated at lower altitudes, and are strongest along the southern
 534 flank of the anticyclone where convective sources are located (Fig. 10e–f). ATAL effects
 535 are of similar magnitude and often exceed the effects of the monsoon anomaly in water
 536 vapor within 370 K–420 K.

537 Overall, the ATAL influence on clear-sky shortwave heating can be thought of as
 538 a direct forcing that compensates for reduced shortwave absorption in the ozone valley
 539 while deepening and expanding the positive effects of water vapor. ATAL effects may
 540 be especially influential on diabatic heating in the western and northern parts of the an-
 541 ticyclone, where the positive effects of water vapor are weak but the negative impacts
 542 of the monsoon ozone valley are relatively strong. These are important regions for mod-
 543 ifying the characteristics of air reaching the stratosphere. In stark contrast to the south-
 544 eastern part of the anticyclone, temperatures in the west are relatively warm and rela-
 545 tive humidities are relatively low owing to adiabatic compression while diabatic mass
 546 transport remains upward (Tegtmeier et al., 2022). Enhanced shortwave heating in these
 547 largely clear-sky regions of the anticyclone (see also Vernier et al., 2015) could poten-

548 tially help air to reach the stratosphere more quickly while avoiding the southeastern ‘cold
 549 trap’; however, the net impact is not so simple to deduce. Additional diabatic heating
 550 can be partitioned in multiple ways. In one limit it can amplify local ascent, moving more
 551 air upward to smaller pressures without changing temperature. This possibility implies
 552 a weakening or reversal of local adiabatic warming and therefore increased humidity, with
 553 potential implications for concentrations of water vapor, short-lived halogenated species,
 554 and other components affected by the formation and sedimentation of ice. In the other
 555 limit it can result in local warming, shifting isentropic surfaces to larger pressures with-
 556 out changing pressure vertical velocities. Such a warming could enhance isentropic as-
 557 cent around the anticyclone, but could also simply be balanced by enhanced longwave
 558 cooling via thermal relaxation either locally or downstream. These possibilities require
 559 evaluation in a comprehensive model framework that can represent coupled interactions
 560 between the ATAL and the anticyclone.

561 4.2 Discrepancies among aerosol reanalysis and forecast products

562 Several other recent reanalyses and forecasts of atmospheric composition are used
 563 to approximate uncertainty windows for ATAL effects on clear-sky shortwave radiative
 564 heating (Table 2). One of these, the Copernicus Atmosphere Monitoring Service (CAMS)
 565 and European Centre for Medium-range Weather Forecasts (ECMWF) Atmospheric Com-
 566 position Reanalysis 4 (CAMS-EAC4; Inness et al., 2019) covers the entire period 2011–
 567 2020 but with a different model and data assimilation system and different emissions re-
 568 lative to MERRA-2 (Text S2 in Supporting Information). Three other products are used
 569 for July–September 2020 only. GEOS-FP is produced using a newer version of the same
 570 atmospheric model and data assimilation system as MERRA-2 (Lucchesi, 2018), but with
 571 a finer horizontal model grid and the inclusion of nitrate and ammonium aerosols. The
 572 CAMS atmospheric composition forecast product (CAMS-FC; Rémy et al., 2019) has
 573 a finer horizontal resolution and additional vertical levels relative to CAMS-EAC4, in-
 574 cludes nitrate and ammonium aerosols where CAMS-EAC4 does not, and implements
 575 a more realistic coupling between sulfur dioxide and sulfate aerosols. Ten-day chemical
 576 forecasts based on the Whole Atmosphere Community Climate Model (WACCM; Get-
 577 telman et al., 2019) with interactive chemistry (Emmons et al., 2020) and aerosols (Liu
 578 et al., 2016) are also considered, with forecast initial conditions taken from GEOS-FP.
 579 Unlike MERRA-2, CAMS-EAC4, and GEOS-FP, AOD observations are not assimilated
 580 in the CAMS-FC or WACCM products used here.

581 These five products show significant differences in their representations of the ATAL
 582 above the monsoon core domain (22.5°N–25°N and 87.5°E–90°E; Fig. 11). GEOS-FP (Fig. 11c)
 583 and WACCM (Fig. 11f) each show ATAL amplitudes approximately twice as large as
 584 the other products, though with substantially different vertical distributions and com-
 585 positions. In WACCM, OC is the largest contributor by mass in the lower part of the
 586 layer ($p > 100$ hPa), switching to sulfate in the upper part of the layer ($p < 100$ hPa).
 587 The maximum in aerosol mass mixing ratio is around 140–160 hPa. In GEOS-FP, con-
 588 centrations of OC, nitrate, ammonium, and sulfate are all similar in magnitude through-
 589 out the depth of the ATAL, with each species (and total mass mixing ratio) peaking
 590 much higher in the layer at ~ 100 hPa. On the other side of the spectrum is the CAMS-
 591 EAC4 reanalysis (Fig. 11d), which produces a weak ATAL mostly composed of OC that
 592 peaks around 150 hPa. Results for CAMS-EAC4 from July–September 2020 are simi-
 593 lar and are not shown here. OC in the CAMS-EAC4 ATAL is mainly hydrophobic in the
 594 lower part of the layer, while all other products indicate that hydrophilic OC outweighs
 595 hydrophobic OC through the full depth of the ATAL. The CAMS-FC product (Fig. 11e)
 596 includes nitrate and ammonium in addition to the species simulated in CAMS-EAC4.
 597 The OC fraction in CAMS-FC is again large in comparison to MERRA-2 and GEOS-
 598 FP, but unlike CAMS-EAC4 is primarily in the hydrophilic component. The distribu-
 599 tion of sulfate is also considerably different, likely due to the revised coupling of SO_4 and
 600 SO_2 (Rémy et al., 2019). This change in the sulfate distribution relative to CAMS-EAC4

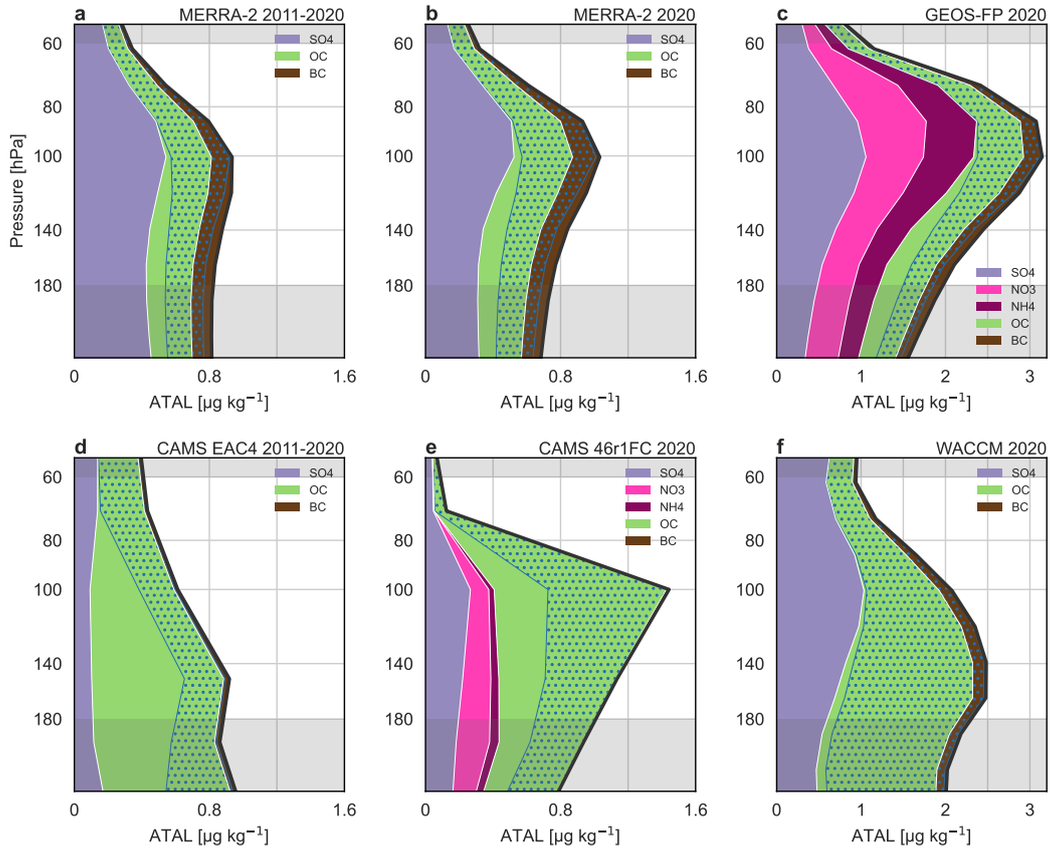


Figure 11. Comparison of ATAL vertical structure and composition in the core analysis region (22.5°N – 25°N , 87.5°E – 90°E) for (a) MERRA-2 during July–September (JAS) 2011–2020, (b) MERRA-2 during JAS 2020, (c) GEOS-FP during JAS 2020, (d) the CAMS reanalysis during JAS 2011–2020, (e) CAMS operational forecasts during JAS 2020, and (f) WACCM hindcasts during JAS 2020. The blue dotted regions mean hydrophilic organic carbon and hydrophilic black carbon. The range of the abscissa is expanded by a factor two in panels (c) and (f) relative to the other four. Among the five products shown, only the GEOS-FP and CAMS forecast products simulated nitrate and ammonium aerosol concentrations.

Table 2. Aerosol reanalysis and forecast products compared in section 4.2. ATAL effects on TOA and surface fluxes are included for reference.

Name	Years	Model	Aerosol species	ATAL effect
MERRA-2 (NASA)	2011–2020	GEOS 5.12.4 0.5°×0.625° 72 levels	BC, OC, dust SO ₄	TOA: 0.04 W m ⁻² SFC: -0.33 W m ⁻²
GEOS-FP (NASA)	2020	GEOS 5.25.1 0.25°×0.3125° 72 levels	BC, OC, dust SO ₄ , NO ₃ , NH ₄	TOA: 0.25 W m ⁻² SFC: -0.84 W m ⁻²
CAMS-EAC4 (ECMWF)	2011–2020	IFS Cy42r1 ~80 km 60 levels	BC, OC, dust SO ₄	TOA: 0.02 W m ⁻² SFC: -0.17 W m ⁻²
CAMS-FC (ECMWF)	2020	IFS Cy46r1 ~40 km 137 levels	BC, OC, dust SO ₄ , NO ₃ , NH ₄	TOA: 0.14 W m ⁻² SFC: -0.22 W m ⁻²
WACCM (NCAR)	2020	WACCM6 0.9°×1.25° 88 levels	BC, OC, dust SO ₄ , NH ₄	TOA: 0.34 W m ⁻² SFC: -0.56 W m ⁻²

601 contributes to raising the peak mass mixing ratio to 100 hPa in CAMS-FC, but most of
602 this increased ATAL height results from changes in OC. Both CAMS products indicate
603 very small fractions of BC in the ATAL. BC loading in the middle and upper troposphere
604 has been reported as overestimated in aerosol analyses that assimilate vertically-integrated
605 AOD (Bozzo et al., 2020); however, it is not clear to what extent this applies to the ATAL,
606 especially given substantial variations in the biomass burning source from year to year.
607 A complementary presentation of differences in ATAL structure and composition is pro-
608 vided in Fig. S2 in Supporting Information.

609 Our intercomparison suggests that MERRA-2 may overestimate the mass fraction
610 of BC within the ATAL but not necessarily the amount, as the latter is comparable to
611 those simulated by GEOS-FP (with assimilation) and WACCM (without). However, the
612 very small concentrations of BC in both CAMS-EAC4 and CAMS-FC are clearly incom-
613 compatible with the relative abundance of BC in MERRA-2, GEOS-FP, and WACCM. Al-
614 though explaining these differences exceeds the scope of this paper, our idealized exper-
615 iments demonstrate significant implications for the magnitude of the ATAL effect on clear-
616 sky shortwave heating (Fig. 7e). Comparing the ATAL from MERRA-2 to that from GEOS-
617 FP suggests that omission of nitrate and ammonium may reduce the ATAL amplitude
618 in MERRA-2 by about half, mainly impacting on the ratio of absorbing aerosol to scatter-
619 ing aerosol. If nitrate and ammonium are added to WASO, both differences are eval-
620 uated within the idealized radiative transfer calculations described in section 3.2. The
621 qualitative distribution of sulfate is consistent across all products except for CAMS-EAC4,
622 which shows little variation in height. Profiles of hydrophobic to hydrophilic ratios in
623 OC and BC are also broadly consistent across all products except for CAMS-EAC4. The
624 MERRA-2 ATAL is therefore not an outlier among these products, which supports our
625 selection of this dataset as a baseline for the idealized calculations introduced above.

626 To clarify how uncertainties in ATAL structure and composition impact radiative
627 heating near the tropopause, we replace the MERRA-2 ATAL in our base case with aerosol
628 profiles from each product listed in Table 2. Other than aerosol loading in the 60–180 hPa
629 layer, atmospheric conditions are identical for all simulations and no aerosols are included
630 either above or below this layer. Figure 12 shows substantial differences in both the mag-
631 nitude and the structure of ATAL effects on clear-sky shortwave heating, with the cor-

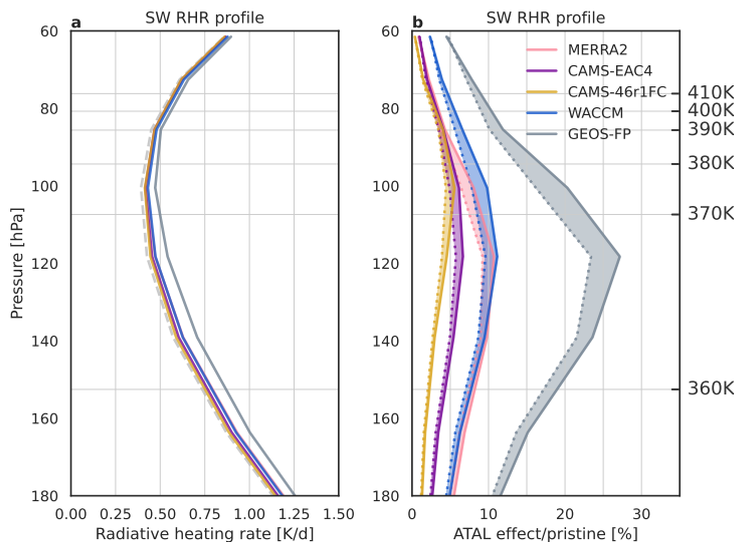


Figure 12. Vertical profiles of (a) clear-sky shortwave radiative heating within the tropopause layer calculated for the no-aerosol case (dashed grey line) and five ATAL profiles (60–180 hPa) from different products and (b) ATAL effects on clear-sky shortwave heating for the different aerosol products relative to the no-aerosol case both with (dotted) and without (solid) adjustment for smaller clear-sky heating rates in libRadtran relative to MERRA-2 (Fig. 5c). All aerosol profiles are time averages for July–September 2020 within the core analysis region (22.5°N–25°N, 87.5°E–90°E).

632 responding impacts on TOA and surface shortwave fluxes listed in Table 2. Consistent
 633 with smaller fractions of absorbing aerosols and weaker ATAL amplitudes, both CAMS-
 634 FC and CAMS-EAC4 produce radiative heating effects that are roughly half the size of
 635 those estimated for MERRA-2. Although the WACCM aerosol layer has approximately
 636 twice as much aerosol by mass as MERRA-2, a larger water soluble fraction and a smaller
 637 amount of hydrophobic black carbon result in similar effects on shortwave heating, with
 638 a smaller peak value but larger increases in heating in the upper part of the layer (Fig. 12b).
 639 The largest radiative effects throughout the layer are produced by GEOS-FP, which sim-
 640 ulates larger fractions of absorbing aerosols in the lower part of the layer and larger con-
 641 centrations of total aerosols in the upper part of the layer (Fig. S2). Aerosol effects on
 642 radiative heating are roughly twice as large when the MERRA-2 ATAL is replaced by
 643 that from GEOS-FP, corresponding to a 15–25% increase relative to clear-sky shortwave
 644 heating rates without aerosol. Accounting for smaller clear-sky shortwave heating based
 645 on libRadtran relative to MERRA-2 (Fig. 5a) reduces the amplitude of the simulated
 646 ATAL effect, but does not change its order of magnitude. Given conservative choices for
 647 optical properties (i.e., assigning the entirety of hydrophilic OC and BC to WASO and
 648 omitting any absorbing component of OC), it is unlikely that this difference leads us to
 649 overestimate the relative ATAL effect on shortwave heating as represented these five aerosol
 650 products. Indeed, interpreting part of the difference between the MERRA-2 and libRad-
 651 tran profiles in Fig. 5a as a deficit in the aerosol contribution rather than the no-aerosol
 652 heating could increase the relative ATAL effect on clear-sky shortwave heating by a com-
 653 parable or even larger amount.

5 Summary and outlook

This research has examined aerosol effects on shortwave fluxes at the surface and nominal top-of-atmosphere (TOA) along with radiative heating near the tropopause above the Asian monsoon. Our analysis confirms that MERRA-2 simulates a well-defined Asian tropopause aerosol layer composed of sulfate, carbonaceous aerosols, and dust (section 3.1). As emissions from volcanic eruptions are omitted after 2010 in MERRA-2 and dust is found to have little impact on radiative heating at these altitudes (Fig. 5a), the ATAL as examined above (sulfate plus carbonaceous aerosols) can be treated as mainly anthropogenic in origin. The ATAL is formed by Asian summer monsoon deep convection, with carbonaceous aerosols contributed mainly by deep convection over the Himalayan-Gangetic Plain and Sichuan Basin as previously reported by W. K. M. Lau et al. (2018) and a substantial convective source of sulfate over southern China (section 3.1). The latter is most influential during the late monsoon season (August–September).

Our simulations clarify ATAL impacts on clear-sky shortwave fluxes at the TOA and surface (section 3.2), with net effects for the time-mean ATAL relative to the no-aerosol case amounting to a 0.04 W m^{-2} reduction of incoming solar radiation at the TOA and a 0.32 W m^{-2} reduction in absorbed shortwave radiation at the surface. These effects increase linearly with increasing total ATAL mass per unit area while holding composition fixed or increasing (decreasing) relative mass fractions of scattering (absorbing) aerosol while holding total ATAL mass fixed. Comparison of these effects as simulated for recent aerosol analysis and forecast products (Table 2) shows large discrepancies in both, with the TOA effect varying by more than an order of magnitude and the surface effect varying by a factor 2–3. Aerosol effects on radiative heating account for around 10% of the clear-sky shortwave heating near the tropopause based on MERRA-2 (section 3.1), with a range of 5–25% calculated for other recent reanalysis and forecast aerosol products (section 4.2). Near the tropopause, ATAL effects based on MERRA-2 are comparable in magnitude to those of monsoon-related anomalies in ozone and water vapor (section 4.1), and are unique among these three in vertical location and horizontal extent. Possible implications for transport to the stratosphere are discussed at the end of section 4.1, but will require evaluation in model systems that represent coupled dynamical interactions between the ATAL and the monsoon anticyclone.

Although the inclusion of interactive aerosol within the meteorological reanalysis framework in MERRA-2 is undeniably helpful, there are large uncertainties that remain to be resolved. In particular, MERRA-2 and other analysis products that assimilate only vertically-integrated AOD may overestimate the abundance of absorbing aerosols at these altitudes (e.g., Bozzo et al., 2020). Assessment of this possibility is complicated by large regional and interannual variations in biomass burning sources and their proximity to convection. Moreover, nitrate aerosols, suggested to be a major component of the ATAL by observations (Höpfner et al., 2019) and some model simulations (Gu et al., 2016), are not represented in MERRA-2. Comparison with GEOS-FP, which uses a newer version of the same model, suggests that excluding nitrate may cause MERRA-2 to underestimate the amplitude of both the ATAL and its effects on radiative heating (section 4.2). However, the separability of absorbing and scattering aerosol contributions to ATAL radiative effects and the largely linear relationships to both components (section 3.2) offer promise for constraining the radiative and dynamical effects of these uncertainties, as they suggest that these effects could be represented within a relatively simple linearized framework. Future work will explore this possibility.

Open Research

MERRA-2 reanalysis products are available through the NASA Goddard Earth Sciences Data Information and Services Center (Global Modeling and Assimilation Office, 2015a, 2015b, 2015d, 2015c). The CAMS-EAC4 reanalysis (Copernicus Atmosphere Monitor-

ing Service, 2020) and CAMS forecast (Copernicus Atmosphere Monitoring Service, 2021) products are available through the Copernicus Atmosphere Data Store. WACCM forecast products are available through NCAR’s Research Data Archive (dataset 313.6; Atmospheric Chemistry Observations & Modeling, National Center for Atmospheric Research, University Corporation for Atmospheric Research, 2020). The libRadtran radiative transfer model is available for download at <http://www.libradtran.org>.

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