Export efficiency of black carbon aerosol in continental outflow: 
Global implications

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[1] We use aircraft observations of Asian outflow from the NASA Transport and Chemical Evolution over the Pacific (TRACE-P) mission over the NW Pacific in March—April 2001 to estimate the export efficiency of black carbon (BC) aerosol during lifting to the free troposphere, as limited by scavenging from the wet processes (warm conveyor belts and convection) associated with this lifting. Our estimate is based on the enhancement ratio of BC relative to CO in Asian outflow observed at different altitudes and is normalized to the enhancement ratio observed in boundary layer outflow (0–1 km). We similarly estimate export efficiencies of sulfur oxides (SOx = SO2(g) + fine SO42− and total inorganic nitrate (HNO3 = HNO3(g) + fine NO3) for comparison to BC. Normalized export efficiencies for BC are 0.63–0.74 at 2–4 km altitude and 0.27–0.38 at 4–6 km. Values at 2–4 km altitude are higher than for SOx (0.48–0.66) and HNO3 (0.29–0.62), implying that BC is scavenged in wet updrafts but not as efficiently as sulfate or nitrate. Simulation of the TRACE-P period with a global three-dimensional model (GEOS-CHEM) indicates that a model timescale of 1 ± 1 days for conversion of fresh hydrophobic to hydrophilic BC provides a successful fit to the export efficiencies observed in TRACE-P. The resulting mean atmospheric lifetime of BC is 5.8 ± 1.8 days, the global burden is 0.11 ± 0.03 Tg C, and the decrease in Arctic snow albedo due to BC deposition is 3.1 ± 2.5%.


1. Introduction

[2] Black carbon (BC), operationally defined as the light-absorbing fraction of carbonaceous aerosols, has complex climatic implications involving atmospheric heating and surface cooling [National Research Council, 2005]. It consists of elemental carbon as well as heavy nonpolar organic compounds [Marley et al., 2001]. It is emitted to the atmosphere by incomplete combustion [Cooke et al., 1999]. The freshly emitted BC is mostly hydrophobic and eventually becomes hydrophilic by oxidation or coating with sulfate and organics [Langner et al., 1992; Parungo et al., 1994; Liousse et al., 1993, 1996]. This conversion is critical for the incorporation of BC in cloud droplets and subsequent removal by rain out. Most global models assume that it takes place on a timescale of 1 day [Cooke and Wilson, 1996; Cooke et al., 1999; Chin et al., 2002; Chung and Seinfeld, 2002; Park et al., 2003]. Cooke and Wilson [1996] first used this timescale on the basis of their comparisons of simulated versus observed BC concentrations in surface air at Amsterdam Island in the Indian Ocean. A global model simulation of aerosol microphysics by Jacobson [2001a] indicates that 50% of BC mass acquires a hydrophilic coating (>20%) within 1 day of emission. The scavenging efficiency of BC has major consequences for climate forcing since outflow from source regions to the free troposphere mainly takes place by wet processes including warm conveyor belts (WCBs) and convection [Thompson et al., 1994; Stohl, 2001; Cooper et al., 2002; Liu et al., 2003]. Once in the free troposphere, BC can be transported on a global scale because precipitation is infrequent.

[3] We present here an assessment of the export efficiency of BC in Asian outflow using aircraft observations over the NW Pacific from the NASA Transport and Chemical Evolution over the Pacific (TRACE-P) mission in
March–April 2001 [Jacob et al., 2003]. The TRACE-P observations provided an extensive characterization of Asian outflow to the Pacific in early spring when this outflow is particularly strong [Liu et al., 2003]. Most of the outflow was driven by midlatitude cyclones, with WCB lifting ahead of the associated cold fronts and boundary layer outflow behind the fronts [Liu et al., 2003]. East Asia is a major source region for BC, contributing ~30% to global anthropogenic BC emissions, mostly from industry, residential coal burning, and biofuel use [Bond et al., 2004]. We compare the observed vertical profiles of BC to those of other soluble and insoluble species measured aboard the aircraft to provide constraints on BC solubility. We then applied the GEOS-CHEM global chemical transport model [Bey et al., 2001] to the TRACE-P observations in order to test the BC scavenging parameterization commonly used in global models and to examine the consequences for the global burden of BC and the deposition of BC to the Arctic. The latter could make an important contribution to climate warming by decreasing snow albedo [Warren and Wiscombe, 1985; Clarke and Noone, 1985; Hansen and Sato, 2001; Hansen and Nazarenko, 2004].

2. Observations

[4] The TRACE-P aircraft mission was conducted offshore of the Asian Pacific Rim during March–April 2001 and focused on chemical characterization of Asian outflow. It used two aircraft, a DC-8 (12-km ceiling) and a P-3B (7-km ceiling), operating out of Hong Kong and Japan. Measurements aboard both aircraft included BC, aerosol sulfate (SO$_4^{2-}$), aerosol nitrate (NO$_3^-$), SO$_2$, gas phase nitric acid (HNO$_3$), and CO, among other species. BC was measured by optical absorption at 565-nm wavelength using a particle/soot absorption photometer (PSAP) (Radiance Research, Seattle, Washington) with correction for artifact absorption from aerosol scattering [Bond et al., 1999]. A mass absorption efficiency of 7 m$^2$ g$^{-1}$ [Clarke et al., 2004] was assumed to convert to mass concentration. This conversion factor was derived from collocated measurements of PSAP absorption and black carbon mass concentrations from the Asian Pacific Regional Aerosol Characterization Experiment (ACE-Asia) aircraft mission conducted over the NW Pacific concurrently with TRACE-P [Huebert et al., 2003]. Intercomparison flight legs during TRACE-P indicated that the aerosol absorption measurements on the DC-8 were systematically higher than on the P-3B and implied unrealistically low aerosol single scattering albedos above 3 km altitude [Moore et al., 2004]. We only use the P-3B data in our analysis.

[5] Fine aerosol concentrations of SO$_4^{2-}$, NO$_3^-$, and Na$^+$ were measured aboard the P-3B using a particle-into-liquid sampler (PILS) with an upper size cutoff of 1.3 μm [Weber et al., 2003]. We use the Na$^+$ data to remove the sea salt contribution to SO$_4^{2-}$ on the basis of the Na$^+/SO_4^{2-}$ ratio in seawater [Keene et al., 1986]. SO$_2$ and HNO$_3$ measurements aboard the P-3B were made using atmospheric pressure ionization mass spectrometry [Tu et al., 2003] and chemical ionization mass spectrometry (CIMS) [Zondlo et al., 2003], respectively. We refer to the sum of SO$_2$ and fine non-sea-salt (nss) SO$_4^{2-}$ as total sulfur oxides (SO$_x$), and we refer to the sum of HNO$_3$ and fine NO$_3^-$ as total inorganic nitrate (HNO$_3^+$). Gas phase sulfuric acid (H$_2$SO$_4$) was also measured on the P-3B by CIMS [Mauldin et al., 2003] but represents <1% of SO$_2$ and was not included in our analysis. Because of the upper size cutoff of PILS the P-3B SO$_2$ and HNO$_3^+$ data did not include a substantial fraction of aerosol sulfate and nitrate associated with coarse soil dust and sea-salt aerosols as indicated by bulk aerosol filter measurements aboard the DC-8 [Jordan et al., 2003]. Measurements of CO were made by tunable diode laser spectrometry (differential absorption CO measurement instrument) [Sachse et al., 1987]. All data used here are averages along the flight tracks on the GEOS-CHEM 2° × 2.5° horizontal grid.

3. Export Efficiencies Constrained by Observations

[6] Koike et al. [2003] previously used the TRACE-P data to estimate export efficiencies of SO$_x$ and total reactive nitrogen oxides (NO$_x$) from east Asia. Similar to BC, SO$_x$ and NO$_x$ originate mainly from combustion and are removed from the atmosphere by deposition. By reference to CO as an inert combustion tracer, Koike et al. [2003] computed the altitude-dependent export efficiencies in east Asian outflow as the ratios of the species enhancements above their background to the corresponding CO enhancements, normalized by the emission ratio for the source region. We can apply this approach to estimate the export efficiency $f_X(z)$ of any combustion-derived species $X$ at altitude $z$:

$$f_X(z) = \frac{1}{R_X} \left( \frac{\Delta X}{\Delta CO} \right)(z),$$

where $R_X$ is the emission ratio of $X$ to CO and $\Delta$ denotes the concentration enhancements relative to background.

[7] We focus on TRACE-P outflow observations north of 30°N in our application of equation (1), as these originated from a well-defined east Asian source region including China, Korea, and Japan [Liu et al., 2003]. Outflow south of 30°N often had a large influence from biomass burning in Southeast Asia with different emission ratios. For comparison of SO$_x$ and BC export efficiencies we further restrict our attention to data west of 140°E because of SO$_2$ emissions from an active volcano on Miyakejima Island (34°N, 139°E) [Koike et al., 2003]. The TRACE-P outflow observations were typically taken 1–3 days downstream from the points of emission in the east Asian source region [Fuelberg et al., 2003; Russo et al., 2003].

[8] Emission ratio estimates for east Asia (defined here as China, Korea, and Japan) are needed for application of equation (1). We use an east Asian BC emission for the 3-month TRACE-P period of 0.45 Tg C, including 0.42 Tg C from anthropogenic sources [Bond et al., 2004] and 0.03 Tg C from biomass burning. The small biomass burning contribution was estimated by applying BC emission factors from Andreae and Merlet [2001] to dry mass burned data with monthly resolution averaged over 1997–2000 from Duncan et al. [2003]. We use a CO emission estimate of 19.2 Tg C for the same region and period as obtained by Palmer et al. [2003] by fitting the TRACE-P CO observations with an inverse model. The resulting BC/CO emission
also compute a normalized export efficiency normalized export efficiency. We propagate into our calculation of the export efficiency. We TRACE-P by a complicated approach involving the CO- background CO for individual aircraft measurements in uncertain [2003] on the basis of the Streets et al. [2003] regional inventory for the TRACE-P period. The CO source for the east Asian region in that inventory is 30% lower than the values (0.069 and 0.070) used by Koike et al. [2003] estimated background concentrations in air upwind from the sources of background is not critical. [11] We also assume background values of 50 ppt for BC and 100 ppt for SO2 and HNO3 as the low tails of the frequency distributions of concentrations in the TRACE-P data. These species have short lifetimes, and hence the specification of background is not critical. [12] Figure 1 (left) shows the mean vertical profiles of export efficiency, and Figure 1 (right) shows normalized export efficiency of BC for the ensemble of observations over the (30°–41°N, 124°–140°E) TRACE-P domain. The profiles were constructed by taking the values of ∆BC and ∆CO for individual observations of ∆CO > 10 ppbv, averaging them over 1-km vertical intervals, and then using these average quantities for input to equation (1). The ∆CO threshold of 10 ppbv is intended to select outflow air masses. Koike et al. [2003] used a higher threshold, ∆CO > 30 ppbv, but we find that this does not affect the results significantly and the lower threshold yields more robust statistics. Our analysis thus includes 102 data points at 0–2 km altitude, 49 at 2–4 km, and 9 above 4 km. [13] We estimate export efficiencies for BC of 0.89–1.06 in the boundary layer (0–2 km), 0.67–0.79 at 2–4 km, and 0.28–0.41 at 4–6 km. The decreasing export efficiencies with altitude indicate wet scavenging of BC during vertical transport. The value of 1.06 at 0–1 km is not significantly different from 1, considering the uncertainty in the approach, in particular the low bias in the BC emission inventory as discussed in section 4. [14] We also show in Figure 1 the export efficiencies of SO2 and HNO3, calculated by applying the same method and with reference to emissions of SO2 and NOx. The export efficiency of HNO3 represents the fraction of emitted NOx that is exported as HNO3. Values for SO2 and HNO3 are 0.25–0.35 and 0.14–0.23 at 0–2 km, 0.17–0.23 and 0.07–0.14 at 2–4 km, and 0.06–0.21 and 0.06–0.12 at 4–6 km, respectively. Koike et al. [2003] previously reported export efficiencies for SO2 and NOx of 0.20–0.45 in the boundary layer (0–2 km) and 0.15 in the free troposphere (2–7 km). Our values for SO2 are consistent; our values for HNO3 are lower than for NOx, as would be expected because of the large contribution of insoluble peroxyacetyl nitrate to the exported NOx. Molar ratios of HNO3/NOx and nssSO42-/SO4 in east Asian anthropogenic plumes at 0–1 km altitude were 0.5 and 0.6, respectively [Koike et al., 2003]. [15] The observed BC export efficiencies are much higher than those of SO2 and HNO3 in the boundary layer outflow.

Figure 1. Mean vertical profiles of (left) export efficiency and (right) normalized export efficiency for black carbon (BC) (squares), SO2, defined as the sum of SO2 and fine non-sea-salt sulfate aerosol (diamonds), and HNO3 T defined as the sum of HNO3 and fine inorganic nitrate aerosol (asterisks). The export efficiencies are derived by application of equations (1) and (2) to the Transport and Chemical Evolution over the Pacific (TRACE-P) P-3B measurements over the NW Pacific (30°–41°N, 124°–140°E). The number of measurements used to calculate export efficiencies at each altitude is also shown.

This normalized export efficiency should overestimate the actual export efficiency because it does not account for scavenging in boundary layer outflow as measured by $f_X(0)$. However, it has the advantage of being independent of $R_X$ as long as one assumes that the same value of $R_X$ applies to outflow at all altitudes.

Application of equations (1) and (2) requires estimates of background concentrations in air upwind from the source region [Mauzerall et al., 1998], especially for CO which has a substantial background because of its long atmospheric lifetime. Koike et al. [2003] estimated background CO for individual aircraft measurements in TRACE-P by a complicated approach involving the CO- CO2 relationship. We use here the inverse model analysis of Palmer et al. [2003] that fitted GEOS-CHEM sources of CO to match the TRACE-P CO observations and decomposed CO concentrations along the TRACE-P flight tracks in terms of contributions from different source regions and source types. We define the local CO background along the aircraft flight tracks as the sum of contributions from the oxidation of methane, oxidation of biogenic nonmethane volatile organic compounds, and anthropogenic sources outside of east Asia. We then add to that value a 16-ppbv enhancement from circumpolar transport of CO emitted by east Asian anthropogenic sources, as determined from the average concentration contributed by these sources in the lower troposphere (0–5 km) over the Middle East (75°E) upwind of east Asia. Our resulting means and standard deviations of background CO concentrations for the TRACE-P data set are 129 ± 11 and 116 ± 10 ppbv in the boundary layer (0–2 km) and the free troposphere (2–6 km), respectively.

Figure 1 (left) shows the mean vertical profiles of export efficiency, and Figure 1 (right) shows normalized export efficiency of BC for the ensemble of observations over the (30°–41°N, 124°–140°E) TRACE-P domain. The profiles were constructed by taking the values of ∆BC and ∆CO for individual observations of ∆CO > 10 ppbv, averaging them over 1-km vertical intervals, and then using these average quantities for input to equation (1). The ∆CO threshold of 10 ppbv is intended to select outflow air masses. Koike et al. [2003] used a higher threshold, ∆CO > 30 ppbv, but we find that this does not affect the results significantly and the lower threshold yields more robust statistics. Our analysis thus includes 102 data points at 0–2 km altitude, 49 at 2–4 km, and 9 above 4 km.

We also show in Figure 1 the export efficiencies of SO2 and HNO3, calculated by applying the same method and with reference to emissions of SO2 and NOx. The export efficiency of HNO3 represents the fraction of emitted NOx that is exported as HNO3. Values for SO2 and HNO3 are 0.25–0.35 and 0.14–0.23 at 0–2 km, 0.17–0.23 and 0.07–0.14 at 2–4 km, and 0.06–0.21 and 0.06–0.12 at 4–6 km, respectively. Koike et al. [2003] previously reported export efficiencies for SO2 and NOx of 0.20–0.45 in the boundary layer (0–2 km) and 0.15 in the free troposphere (2–7 km). Our values for SO2 are consistent; our values for HNO3 are lower than for NOx, as would be expected because of the large contribution of insoluble peroxyacetyl nitrate to the exported NOx. Molar ratios of HNO3/NOx and nssSO42-/SO4 in east Asian anthropogenic plumes at 0–1 km altitude were 0.5 and 0.6, respectively [Koike et al., 2003].

The observed BC export efficiencies are much higher than those of SO2 and HNO3 in the boundary layer outflow.
This could be explained by inefficient removal of fresh BC in the source region. In addition, boundary layer outflow is a dry process, and BC (unlike the gas phase species SO\textsubscript{2} and HNO\textsubscript{3}) is not removed efficiently by dry deposition. The uncertainty in BC emission is a complicating factor, as regional modeling by Carmichael et al. [2003] of the BC observations in TRACE-P implies that east Asian BC sources in the Bond et al. [2004] inventory are underestimated. We address that issue further in section 4.

16 The normalized export efficiencies for BC, SO\textsubscript{2}, and HNO\textsubscript{3} in Figure 1 (right) show a decrease with altitude. In the lower free troposphere (2–4 km), which was a prevailing WCB outflow altitude during TRACE-P [Carmichael et al., 2003; Liu et al., 2003], the BC normalized export efficiency (0.63–0.74) is higher than that of SO\textsubscript{2} (0.48–0.66) or HNO\textsubscript{3} (0.29–0.62). The higher normalized export efficiency for BC relative to SO\textsubscript{2} and HNO\textsubscript{3} implies some resistance to scavenging and thus some hydrophobic character. At 5–6 km, there is no significant difference in the normalized export efficiencies of HNO\textsubscript{3}, SO\textsubscript{2}, and BC (0.17–0.27). This could reflect a more thorough processing of the outflow by precipitation, but there are only nine data points contributing to the calculation of export efficiencies above 4 km, and thus the estimates are uncertain.

4. Model Simulation: Implications for Black Carbon Scavenging

17 We use the GEOS-CHEM three-dimensional model of aerosol-oxidant chemistry [Bey et al., 2001; Park et al., 2003, 2004] to examine the constraints from the TRACE-P observations on the standard representation of BC scavenging used in global models. The GEOS-CHEM model (version 6.02, available at http://www-as.harvard.edu/chemistry/trop/geo) uses assimilated meteorological data for the TRACE-P period from the NASA Goddard Earth Observing System (GEOS-3) including winds, convective mass fluxes, mixing depths, temperature, precipitation, and surface properties. The data have 6-hour temporal resolution (3-hour for surface variables and mixing depths), 1\textdegree\times1\textdegree horizontal resolution, and 48 sigma vertical layers. We degrade the horizontal resolution to 2\textdegree\times2.5\textdegree for input to GEOS-CHEM.

18 The simulation of BC aerosols in GEOS-CHEM is as described by Park et al. [2003] with updated emissions as described below. The simulation of SO\textsubscript{2} and HNO\textsubscript{3} species is as described by Park et al. [2004] with nitrate aerosol formation driven by H\textsubscript{2}SO\textsubscript{4}-HNO\textsubscript{3}-NH\textsubscript{3}-H\textsubscript{2}O thermodynamics. Wet deposition follows the scheme of Liu et al. [2001], including contributions from scavenging in convective updrafts, rain out from convective anvils, rain out and washout from large-scale precipitation, and species-dependent release of gases upon cloud freezing [Mari et al., 2000]. The scheme was originally tested by simulation of aerosol radionuclides [Liu et al., 2001] and also yields a good simulation of soluble species concentrations in the United States [Park et al., 2003, 2004] and in North American outflow [Li et al., 2004]. Dry deposition is simulated with a standard resistance-in-series model dependent on local surface type and meteorological conditions, as described by Wang et al. [1998]; it is important for gaseous SO\textsubscript{2} and HNO\textsubscript{3} but not for fine aerosols.

19 The model simulation of BC treats hydrophobic and hydrophilic BC as two separate transported species. Following the standard parameterization used in global models, we assume that all BC is emitted as hydrophobic and becomes hydrophilic with an e-folding time \(\tau\). Global models of BC commonly assume \(\tau = 1\) day, as mentioned previously (in section 1). We examine here the constraints on \(\tau\) from the TRACE-P observations. Wet deposition is applied only to hydrophilic BC. Hydrophobic BC can also be scavenged by collision with cloud droplets and falling raindrops, but the effect on aerosol mass is small [Jacobson, 2003].

20 We use global anthropogenic (fuel) emissions of BC (4.8 Tg C yr\textsuperscript{-1}) from the gridded annual Bond et al. [2004] inventory for 1996. The BC emissions from fossil fuel and biofuel in east Asia are 1.2 and 0.43 Tg C yr\textsuperscript{-1}, respectively. Biomass burning emissions are specified from a gridded climatological inventory with monthly resolution [Duncan et al., 2003] and using BC emission factors from Andreae and Merlet [2001]. The resulting global annual BC emission from biomass burning is 2.4 Tg C yr\textsuperscript{-1}. Biomass burning emissions in east Asia during TRACE-P are 0.03 Tg C, much lower than fuel emissions.

21 Model simulations are conducted from August 2000 to December 2001. The first 5 months are used for initialization. We focus on results for 2001. For comparison with the TRACE-P observations we sample the 3-hour average model results along the TRACE-P flight tracks and for the flight periods.

22 Figure 2 compares mean observed and simulated vertical concentration profiles of SO\textsubscript{2}, HNO\textsubscript{3}, and fine NO\textsubscript{3}\textsuperscript{-} for the ensemble of TRACE-P flights over the east Asian outflow domain (30\degree–41\degree N, 124\degree–140\degree E). The simulated and observed SO\textsubscript{2} concentrations decrease rapidly with altitude, and there is no significant model bias. The simulated HNO\textsubscript{3} concentration is a factor of 1.8 higher than observed. An explanation is that the model does not account for the uptake of HNO\textsubscript{3} on coarse soil dust and sea-salt aerosols [Grassian, 2001] which would be excluded from the HNO\textsubscript{3} measurement. Bulk aerosol measurements from the DC-8 aircraft during TRACE-P indicated that 50% of total inorganic nitrate and 20% of total sulfur oxides, on average, were incorporated in coarse soil dust aerosols in dust-impacted Asian outflow [Jordan et al., 2003]. Recent measurements in Japan during the Asian dust season [Ooki and Uematsu, 2005] indicated that 50% of HNO\textsubscript{3} reacted with soil dust aerosols, on average, but only 10% of SO\textsubscript{2} reacted. Aerosol nitrate in the model is mainly fine NH\textsubscript{4}NO\textsubscript{3} and reproduces well the fine NO\textsubscript{3}\textsuperscript{-} observations aboard the P-3B, as shown in Figure 2. Overall, the good simulation of SO\textsubscript{2} and fine NO\textsubscript{3}\textsuperscript{-} indicates a successful simulation of Asian outflow and associated scavenging for the TRACE-P period.

23 We also compare in Figure 2 the mean observed and simulated vertical concentration profiles of BC, using \(\tau = 1\) day in the model to describe BC scavenging. The observed BC concentration is, on average, 60% higher than the simulation, which could reflect either insufficient emission or excessive scavenging. We test for the latter in Figure 3 by comparing simulated and observed values of the normalized export efficiencies for BC. For this comparison we use TRACE-P outflow data extending east to 150\degree E (30\degree–
41°N, 124°–150°E) to have better statistics since comparison to SOx export efficiency is not an issue. The change in domain results in a slight increase (≈5%) in the computed BC export efficiency relative to that in Figure 1. The normalized export efficiencies in the model are computed in the same way as in the observations and for model values of \( t \) ranging from 0 to 5 days to test the sensitivity to \( t \).

From Figure 3 we see that \( t = 1 \) day (as assumed in our standard simulation) reproduces the observed vertical profile of the normalized export efficiency, implying that scavenging bias is not responsible for the model underestimate of BC concentrations in Figure 2. Emissions in the Bond et al. [2004] inventory are likely too low, as suggested also in previous analyses of the TRACE-P data [Carmichael et al., 2003; Clarke et al., 2004]. Test simulations indicate that increasing east Asian emissions by 60% relative to the Bond et al. [2004] inventory would correct the discrepancy in the GEOS-CHEM simulation. Increasing \( t \) up to 5 days would not.

5. Implications for the Global BC Budget and for Deposition to the Arctic

Comparison of model results to TRACE-P observations in Figure 3 shows that our best estimate of the timescale \( t \) for conversion of hydrophobic to hydrophilic BC in global models is 1 day, with a likely range of 0–2 days (i.e., \( t = 1 \pm 1 \) days). We examine the implications for the simulation of the global burden of BC and the deposition flux to the Arctic north of 70°N. Figure 4 shows the simulated BC burden, annual wet deposition flux in the Arctic, and corresponding decrease in Arctic snow albedo as a function of \( t \). The global burden of BC increases by a factor of 3 as \( t \) increases from 0 to 5 days. A value for \( t \) of 1 ± 1 days combined with the Bond et al. [2004] global inventory yields a global BC burden of 0.11 ± 0.03 Tg C, lower than values of 0.17–0.32 Tg C found in previous global model studies [Cooke and Wilson, 1996; Jacobson, 2001b; Chuang et al., 2002; Chung and Seinfeld, 2002; Chin et al., 2002; Wang, 2004]. This reflects our lower BC emission (7.2 Tg C yr\(^{-1}\)) than was used in those previous studies (12–18.7 Tg C yr\(^{-1}\)). An Intergovernmental Panel on Climate Change [2001] intercomparison of global model simulations with imposed BC emission of 12.4 Tg C yr\(^{-1}\) found a mean high bias of a factor of 2.3 compared to observations averaged over all sites, although with considerable scatter. The Bond et al. [2004] inventory would partly correct this global mean bias, although as we have seen, it is too low in east Asia. With \( t = 1 \pm 1 \) days we compute in the
model a mean atmospheric lifetime for BC of 5.8 ± 1.8 days, which is within the range of 4.0–7.9 days found in the above-mentioned studies.

[25] Our simulated annual BC deposition flux of 50–70 Gt C yr⁻¹, with a time constant of 30–190 days, is within the range of 58 ± 12 Gt C yr⁻¹ found in observations [Clarke and Noone, 1985]. Our simulated BC scavenging rate of 1 ± 1 days is 58 ± 12 Gt C yr⁻¹.

[26] Although our TRACE-P simulation shows that a model time constant τ = 1 ± 1 days for conversion of emitted BC from hydrophobic to hydrophilic can fit the TRACE-P observational constraints on export efficiencies in Asian outflow, the crudeness of this parameterization must be acknowledged. The value of τ should vary depending on the local environment, and it is doubtful that a single time constant can properly describe the aging of BC and the implications for its scavenging efficiency. Further work is necessary to examine this aging and scavenging, both through observations and through improved models.

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Figure 2. Mean vertical profiles of simulated (solid circles) versus observed (squares) concentrations of SO\(_x\) (red), HNO\(_3\) (green), fine NO\(_3^-\) (blue), and BC (black) for the ensemble of TRACE-P P-3B observations over the NW Pacific (30°–41°N, 124°–140°E). The model results were sampled along the aircraft flight tracks. The data were binned in 1-km vertical intervals and were then averaged to construct the profiles.

Figure 3. Comparison of simulated (solid circles) versus observed (squares) normalized export efficiencies of BC for the ensemble of observations and for model values of \(\tau\) ranging from 0 to 5 days as indicated by the inset. The normalized export efficiencies are calculated using equation (2) for the ensemble of TRACE-P P-3B flights over the domain (30°–41°N, 124°–150°E). The number of measurements used to calculate export efficiencies at each altitude is also shown.