# Correction published 23 July 2005

# Control of fossil-fuel particulate black carbon and organic matter, possibly the most effective method of slowing global warming

Mark Z. Jacobson

Department of Civil and Environmental Engineering, Stanford University, Stanford, California, USA

Received 9 October 2001; revised 5 February 2002; accepted 12 April 2002; published 15 October 2002.

[1] Under the 1997 Kyoto Protocol, no control of black carbon (BC) was considered. Here, it is found, through simulations in which 12 identifiable effects of aerosol particles on climate are treated, that any emission reduction of fossil-fuel (f.f.) particulate BC plus associated organic matter (OM) may slow global warming more than may any emission reduction of CO<sub>2</sub> or CH<sub>4</sub> for a specific period. When all f.f. BC + OM and anthropogenic  $CO_2$  and  $CH_4$  emissions are eliminated together, the period is 25–100 years. It is also estimated that historical net global warming can be attributed roughly to greenhouse gas plus f.f. BC + OM warming minus substantial cooling by other particles. Eliminating all f.f. BC + OM could eliminate 20–45% of net warming (8–18% of total warming before cooling is subtracted out) within 3-5 years if no other change occurred. Reducing CO<sub>2</sub> emissions by a third would have the same effect, but after 50-200 years. Finally, diesel cars emitting continuously under the most recent U.S. and E.U. particulate standards (0.08 g/mi; 0.05 g/km) may warm climate per distance driven over the next 100+ years more than equivalent gasoline cars. Thus, fuel and carbon tax laws that favor diesel appear to promote global warming. Toughening vehicle particulate emission standards by a factor of 8 (0.01 g/ mi; 0.006 g/km) does not change this conclusion, although it shortens the period over which diesel cars warm to 13–54 years. Although control of BC + OM can slow warming, control of greenhouse gases is necessary to stop warming. Reducing BC + OM will not only slow global warming but also improve human health. INDEX TERMS: 0305 Atmospheric Composition and Structure: Aerosols and particles (0345, 4801); 0320 Atmospheric Composition and Structure: Cloud physics and chemistry; 0345 Atmospheric Composition and Structure: Pollution-urban and regional (0305); 1620 Global Change: Climate dynamics (3309)

Citation: Jacobson, M. Z., Control of fossil-fuel particulate black carbon and organic matter, possibly the most effective method of slowing global warming, *J. Geophys. Res.*, 107(D19), 4410, doi:10.1029/2001JD001376, 2002.

#### 1. Introduction

[2] To date, several studies have advocated non-CO<sub>2</sub> greenhouse-gas emission controls in conjunction with CO<sub>2</sub> emission controls as a method of slowing global warming [e.g., Hayhoe et al., 1999; Hansen et al., 2000; Smith et al., 2000; IPCC, 2001]. With respect to particulates, early global radiative studies of black carbon (BC) treated it as either externally mixed or well mixed internally [e.g., Haywood et al., 1997]. Jacobson [2000] argued that treating BC as well mixed internally was not physical and treating it as externally mixed was not representative of the real mixing state of BC. It was hypothesized that a more representative optical treatment of BC might be that of a core surrounded by shell. Under that assumption, the global direct forcing of fossil fuel plus biomass-burning BC was 0.5 W/m<sup>2</sup>, implying that BC control might slow global warming to some degree. This implication was subsequently supported by Hansen et al. [2000] and Smith et al. [2000]. An unstudied issue, though, was the actual

- mixing state of BC. *Jacobson* [2001a] performed simulations of the evolution of the global scale mixing state and direct forcing of BC. It was found that both appeared to approach those of an internal mixture with a core rather than those of an external mixture, supporting the contention that BC has a strong direct forcing and its control might slow global warming.
- [3] Among constituents causing warming, anthropogenic CO<sub>2</sub>, BC, and CH<sub>4</sub> may have the greatest direct forcing, although other gases and particles also contribute to global warming. Other warming gases include tropospheric O<sub>3</sub> (whose production also leads to the formation of particulate nitrate and secondary organics, which enhance cooling), halocarbons, and N<sub>2</sub>O. Other warming particle components include iron, aluminum, ammonium, polycyclic aromatics, and nitrated aromatics [*Jacobson*, 2001b]. To date, one study has examined the short-term climate response to size-resolved aerosols containing BC as a core on the urban scale [*Jacobson*, 1997a, 1997b], but no study has compared the effects over time of reducing BC, CO<sub>2</sub>, and CH<sub>4</sub> emissions on the global scale.
- [4] To estimate the relative global-scale climate benefits of BC, CO<sub>2</sub>, and CH<sub>4</sub> emission controls, time-dependent

global model simulations were carried out. The model used in this study is described in section 2. During the simulations, aerosol particles fed back to temperatures in at least 12 ways. These are described in section 3. Subsequent sections of this paper describe the simulations (section 4), comparisons of baseline predictions with observations (section 5), climate response results (section 6), and a comparison of the effects of diesel and gasoline on global warming (section 7).

# 2. Description of the Model

[5] The model used was GATOR-GCMM (Gas, Aerosol, Transport, Radiation, General Circulation, and Mesoscale Meteorological model) [*Jacobson*, 1994, 1997a, 1997b, 2001c, 2001d]. The model treated time-dependent gas, aerosol, cloud, radiative, dynamical, and transport processes over a 4°S-N × 5°W-E grid with 39 sigma-pressure layers (23 tropospheric, 16 stratospheric; 3 in the bottom 1 km) from the ground to 0.425 mb (≈55 km).

#### 2.1. Gas Processes

- [6] Gas processes included emissions, photochemistry, transport (section 2.3), homogeneous nucleation (section 2.2), condensational growth (section 2.2), dissolutional growth (section 2.2), dry deposition, and washout (section 2.5).
- [7] Monthly gridded emission rates of CO<sub>2</sub>, CO, CH<sub>4</sub>, NO<sub>x</sub>, SO<sub>2</sub>, ethene, propene, ethane, and propane from biomass burning were obtained by multiplying a mean biomass-burning emission factor for each gas [Ferek et al., 1998] with the monthly gridded BC biomass-burning emission inventory of Cooke and Wilson [1996], then dividing by a mean emission factor of BC [Ferek et al., 1998]. The BC inventory for biomass burning from Cooke and Wilson was designed for the mid-1980s and gives a BC emission rate of 6 Tg C/yr. Andreae and Merlet [2002] calculated a BC emission rate from biomass burning of 4.7 Tg C/yr using emission factors derived therein plus estimates of biomass burned given by Lobert et al. [1999], which were based on data from 1980 to the early 1990s. To account for the more recent work, the gas inventory for biomass burning was reduced by 25%.
- [8] Other emitted gases included volcanic SO<sub>x</sub> [Andres and Kasgnoc, 1998], NO<sub>x</sub> and other SO<sub>x</sub> [Voldner et al., 1999], NH<sub>3</sub> [Bouwman et al., 1997], CH<sub>4</sub> [Stern and Kaufman, 1998], CO<sub>2</sub> [Marland et al., 1999], and CO, N<sub>2</sub>O, and NMVOCs [Olivier et al., 1996]. Ocean emission rates of DMS were calculated with the parameterization of Liss and Merlivat [1986] and seawater DMS concentrations of Kettle et al. [1999].  $NO_x$  from intercloud lightning was determined by integrating the number of bounceoffs following collisions among size-resolved ice crystals and graupel, calculating an intracloud flash rate from bounceoffs using an equation of Wang and Prinn [2000], assuming the cloud-ground flash rate was 35% of the cloud-cloud flash rate [Boccippio et al., 2001], and assuming  $1.9 \times 10^{26}$  NO molecules per cloud-cloud flash and  $1.9 \times 10^{27} \text{ NO}$ molecules per cloud-ground flash (average of high and low values for each by Wang and Prinn [2000]).
- [9] Gas photochemistry was solved among 40 gases and 109 reactions with SMVGEAR II [Jacobson, 1998a]. Gas

dry deposition velocities were calculated with a threeresistance approach. The surface resistance [Wesely, 1989; Walmsley and Wesely, 1996] depended on landcover type, determined at 1 km resolution from USGS [1999].

#### 2.2. Aerosol and Gas-Aerosol Processes

- [10] The model treated one aerosol size distribution,  $0.01-110~\mu m$  diameter, with 17 moving center [*Jacobson*, 1997a] size bins in each grid cell. The number concentration of total particles and the mole concentrations of  $H_2O(aq)$ ,  $H^+$ ,  $NH_4^+$ ,  $Na^+$ ,  $H_2SO_4(aq)$ ,  $HSO_4^-$ ,  $SO_4^{2-}$ ,  $NO_3^-$ ,  $C1^-$ , BC, organic matter [OM = organic carbon (OC) + functional groups], and soil-dust were prognostic variables in each bin. Particle emission sources included biomass burning, seaspray uplift, soil uplift, and f.f. combustion.
- [11] Emitted biomass-burning components included BC, OM, K<sup>+</sup>, Na<sup>+</sup>, Ca<sup>2+</sup>, Mg<sup>2+</sup>, NH<sub>4</sub><sup>+</sup>, Cl<sup>-</sup>, SO<sub>4</sub><sup>2-</sup>, and NO<sub>3</sub><sup>-</sup>. Since K<sup>+</sup>, Ca<sup>2+</sup>, and Mg<sup>2+</sup> were not carried in the simulations, their mole-equivalent emissions were added to those of Na<sup>+</sup>. Monthly gridded emission rates of each species were obtained in the same manner as were biomass-burning gases (section 2.1). Particulate emissions from biomass burning were distributed lognormally over model size bins [*Reid and Hobbs*, 1998].
- [12] Emitted sea-spray and spume-drop components included  $H_2O$ ,  $K^+$ ,  $Na^+$ ,  $Ca^{2+}$ ,  $Mg^{2+}$ ,  $Cl^-$ , and  $SO_4^{2-}$ . Again, since  $K^+$ ,  $Ca^{2+}$ , and  $Mg^{2+}$  were not carried in the model, their mole-equivalent emissions were added to those of  $Na^+$ . Sea-spray and spume-drop emissions versus size were calculated as a function of wind speed from the works of *Monahan et al.* [1986] and Wu [1993], respectively. Emitted drop composition was set to that of seawater.
- [13] Due to the computational burden, the only emitted soil component in the model was an unreactive, generic soil component. Soil emissions versus size were calculated from the work of *Marticorena et al.* [1997] using soil distribution data from *FAO* [1995]. The soil emission rate depended on wind speed, soil type, and particle size.
- [14] Emitted f.f. particulate components included BC + OM. The f.f. BC inventory was obtained from the work of Cooke et al. [1999] and applied over a lognormal combustion distribution for f.f. [Maricq, 1999a]. Although the inventory of Cooke et al. was designed for the 1980s and may have overestimated powerplant emission factors, Streets et al. [2001] found that the overall current inventory of BC from China was not too different from that of Cooke et al. because Cooke et al. did not include residential combustion of biofuel emission sources of BC. Table 1 here shows that the model underpredicted BC in urban and many rural areas, suggesting that the BC inventory did not lead to overestimates of BC climate effects. Baseline OM/ BC emission ratios from f.f. were set to 3.1:1 (an average OC/EC f.f. ratio from the work of Strader et al. [1999] is 2.4, and the OM/OC ratio was 1.3:1). The OM/BC emission ratio for biomass burning was 8:1.
- [15] H<sub>2</sub>SO<sub>4</sub>–H<sub>2</sub>O homogenous nucleation was calculated with classical theory [*Hamill et al.*, 1982; *Zhao and Turco*, 1995] and solved simultaneously with condensation of H<sub>2</sub>SO<sub>4</sub>–H<sub>2</sub>O between the gas phase and all 17 bins with a mass-conserving, noniterative, and unconditionally stable condensation scheme [*Jacobson*, 1997c]. In addition, dissolution of each NH<sub>3</sub>, HNO<sub>3</sub>, and HCl was solved between

Table 1. Comparison of Modeled Versus Observed Near-Surface BC Concentrations

Station	Latitude	Longitude	Modeled BC (ng/m <sup>3</sup> )	Measured BC (ng/m <sup>3</sup> )	Period	Data reference
				Marine		
Amsterdam Island	37°30′S	77°10′E	8.5	5.5	Annual	Cachier et al. [1994]
Atlantic Ocean	$31-37^{\circ}N$	66-76°W	160	10	January-February	Quinn et al. [2001]
	15-31°N	43-66°W	45	10		
	8-15°N	33-43°W	65	30		
	3-8°N	26-33°W	220	360		
	5°S-3°N	17–26°W	180	$380 \pm 210$		
	25-5°S	6°E-17°W	50	$20 \pm 30$		
	33-24°S	18-6°E	90	10		TT 100 1 5100 67
Bermuda	32°45′N	65°W	70	30	February	Wolff et al. [1986]
		0	80	40	August	
Böisto Island	60°20′N	26°30′E	1030	50-1000	June	Raunemaa et al. [1993]
Chichi-jima	27°N	142°E	175	610	December	Ohta and Okita [1984]
Corsica	42°N	9°E	420	380	Spring	Cachier et al. [1989]
East China Sea	27-32°N	127-129°E	370	355	April-May	Parungo et al. [1994]
	27-33°N	124-129°E	810	505	October-November	
	28-32°N	125-128°E	920	133	December	
Hachi-jima	33°5′N	139°45′E	885	1070	January	Ohta and Okita [1984]
			845	740	December	
Halley Bay	75°24′S	27°W	2.7	0.3 - 3		Cachier et al. [1986],
						Hansen et al. [1988]
La Réunion Island	21°30′S	55°30′E	70	50 - 650	Annual	Bhugwant et al. [2000]
Mace Head, Ireland	53°20′N	9°54′W	490	26 (17-34)	July-August	Krivacsy et al. [2001]
			450	47 - 1041	Annual	Derwent et al. [2001]
NE Atlantic			80	268	October-November	O'Dowd et al. [1993]
NE Pacific	$27-48^{\circ}N$	175°E	60	150 - 300	April-October	Kaneyasu and Murayami [2000
Ny-Alesund	78°54′N	11°53′E	150	70(28-174)	March-May	Heintzenberg [1982]
			170	293 (123-567)	March-April	Clarke [1989]
Oki Island	36°9′N	133°12′W	55	520	Annual	Mukai et al. [1990]
San Nicolas Island	33°9′N	119°18′W	240	260	Annual	Kim et al. [2000]
				Rural		
Abastumani	41°30′N	42°40′E	460	980	Annual	Dzubay et al. [1984]
Abisko	68°18′N	18°30′E	300	393	March-April	Noone and Clarke [1988]
TOISKO	00 1010	10 JUL	360	259-780	Annual	Clarke [1989]
Allegheny	40°N	79°W	820	1300	August	Keeler et al. [1990]
Aspyreten, Sweden	58°48′N	17°23′W	115	100 (50–170)	June-July	Zappoli et al. [1990]
Cheboque Point	43°48′N	66°6′W	270	81-139	•	
*	79°33′N	90°37′E	125		August-September	Chylek et al. [1996]
East Arctic	2°S	77.18°W	265	25-115 $100-520$	March-April	Polissar [1993]
Ecuador	2°15′S	77.18 W 79°52′W	430	100-320 $100-520$	Annual	Andreae et al. [1984]
Florido	28°N	82°W	690		June	Andreas et al [1004]
Florida				400	Annual	Andreae et al. [1984]
Gif sur Yvette	48°42′N	2°8′E	1300	1650 - 2750	Seasonal	Cachier et al. [1989];
	4.600.0/2.1	5050/F	710	200 (0. 720)	T.1. A.	Brémond et al. [1989]
Jungfraujoch, Switzerland		7°59′E	710	290 (8-720)	July-August	Krivacsy et al. [2001]
K-Puszta, Hungary	46°57′N	19°42′E	1670	810	Annual	Heintzenberg and Mészáros [198
			1700	600 (400–600)	July-August	Zappoli et al. [1999]
	4.403.7	4.0***	1700	750 (120–1890)	July-August	Krivacsy et al. [2001]
Landes forest	44°N	1°W	350	300	Autumn	Cachier et al. [1989]
Lin'an	30°15′N	119°42′E	2150	2070	July-August	Parungo et al. [1994]
North Carolina	35°18′N	80°W	550	520	Annual	Andreae et al. [1984]
Nylsvley, South Africa	24°39′N	28°24′E	340	850 (450–1340)	May	Puxbaum et al. [2000]
Petten	52°54′N	2°55′E	1360	1630	April	Berner et al. [1996]
Rorvik	57°18′N	12°12′E	1010	725	January-May	Broström-Lundén et al. [1994]
Senonche	48°34′N	12°E	1100	1200	March	Cachier et al. [1990]
Southwest United States	34°18′N	106°W	160	150 - 220	Annual	Andreae et al. [1984]
West Mountain	40°N	116°E	720	1710 - 2610	August	Parungo et al. [1994]
Maldives	4°58′N	73°28′E	290	2700 - 9400	February	Chowdhury et al. [2001]
				Urban		
Arnhem	52°N	5°54′E	1600	2950	October-November	Janssen et al. [1997]
Atlanta, Ga.	33°N	85°W	460	1500 (500-3000)	August	Modey et al. [2001]
Baltimore	JJ 14	05 <b>**</b>	800	530-2600	July	Brunciak et al. [2001]
Beijing	40°N	116°E	680	8700-10,100	Annual	He et al. [2001]
Denjing	TU 1N	110 E	700	6270	Summer	11e et at. [2001]
			640	10,230	Autumn	
			690	11,080	Winter	
71.:/T -1 3.6° 1.°	41001/27	07020/337	680	6670	Spring	000-1
Chicago/Lake Michigan	41°31′N	87°39′W	970	340-420	January	Offenberg and Baker [2000]
OI 1 D	200465-	7.000 5/22	1050	350-490	July	D
Chesapeake Bay	38°40′N	76°25′W	790	400-7300	July	Brunciak et al. [2001]
Clermont Ferrand	45°46′N	34°E	1270	2400	April	Lyubovtseva and Yatskovich [198

Table 1. (continued)

Station	Latitude	Longitude	Modeled BC (ng/m <sup>3</sup> )	Measured BC (ng/m <sup>3</sup> )	Period	Data reference
Dushanbe	38°33′N	68°48′E	290	10,700-12,000	August-September	Hansen et al. [1993]
Gorlitz	51°10′N	14°59′E	1950	1300	Annual	Zier [1991]
Halle	51°29′N	12°E	1800	1600	Annual	Zier [1991]
Hamburg	53°52′N	9°59′E	2800	1975	Annual	Heintzenberg and Mészáros [1985]
Kaohsiung			1400	3300 - 8000	November-April	Lin and Tai [2001]
Kap Arkona	57°37′N	13°21′E	810	400	Annual	Zier [1991]
Leeds	53°48′N	1°34′W	1020	780 - 1300	Summer, Winter	Willison et al. [1985]
Los Angeles Basin	34°4′N	118°15′W	390	2750 - 4230	Annual	Kim et al. [2000]
Moscow	55°45′N	37°35′E	1260	3500 - 6750	Summer, Winter	Kopeykin et al. [1993]
Mount Gibbes	35°78′N	82°29′E	240	74	Winter	<i>Im et al.</i> [2001]
			200	230	Summer	
Nagoya	35°10′N	136°50′E	800	13,000	Annual	Kadowaki [1990]
Orleans	47°54′N	15°2′E	1180	2900	March	Del Delumyea and Kalivretenos [1987]
Paris	48°50′N	2°20′E	1340	4600	Annual	Ruellan and Cachier [2001]
			1600	13,600	Summer-Autumn	
Potsdam	52°3′N	13°4′E	1950	1000	Annual	Zier [1991]
Po Valley, Italy	44°39′N	11°37′E	800	1000 (500-1500)	September	Zappoli et al. [1999]
Radebeul	51°6′N	13°55′E	2200	1400	Annual	Zier [1991]
Santiago	33°33′S	70°36′W	90	30,600	June	<i>Didyk et al.</i> [2000]
Seoul	37°20′N	126°35′E	2340	8400	June	Park et al. [2001]
Thessaloniki	40°31′N	22°58′E	1080	3500 - 8900	June	Chazette and Liousse [2001]
Uji	34°32′N	135°29′E	860	5210	Winter	Hitzenberger and Tohno [2001]
			680	2600	Summer	
			770	4890	Annual	
Vienna	48°12′N	16°22′E	1370	6580	Winter	Hitzenberger and Tohno [2001]
			1890	2600	Summer	
			1550	4890	Annual	

the gas phase and 17 particle size bins with an analogous dissolution scheme [*Jacobson*, 1997c]. SO<sub>2</sub> dissolution and aqueous chemistry in aerosol particles and cloud drops were treated as described in section 2.5.

[16] Aerosol liquid water content, pH, and ion distributions in all size bins were solved with EQUISOLV II [Jacobson, 1999b]. Aerosol-aerosol coagulation (assuming kernels for Brownian motion, Brownian diffusion enhancement, gravitational collection, turbulent inertial motion, and turbulent shear) was solved among all components and total particles in each bin with a volume-conserving, noniterative, scheme [Jacobson et al., 1994]. Aerosol-cloud coagulation, aerosol rainout, and aerosol washout are discussed in section 2.5. Particle sedimentation and dry deposition were solved as described by Jacobson [1999a, Chapter 20].

### 2.3. Dynamics and Transport

[17] The global dynamics module integrates equations for momentum (under the hydrostatic assumption), thermodynamic energy, and total water. The solution is a fourth-order scheme that conserves potential enstrophy and energy [Arakawa and Lamb, 1981]. Transport of gases and aerosols was carried out with the scheme of Walcek and Aleksic [1998] using online winds predicted by the dynamics module and vertical diffusion coefficients predicted by the turbulence module.

# 2.4. Turbulence, Stratus Clouds, Cumulus Clouds, and Species Transport in Clouds

[18] The turbulence closure (order 2.5) and stratus cloud parameterizations in the model were coupled together, and both were drawn from the work of *Mellor and Yamada* [1982]. Turbulence parameters affected momentum, energy,

and tracers. The stratus cloud scheme predicted stratus cloud fraction and water content in each layer given turbulence terms and vertical gradients of energy and moisture. Cloud water and fraction fed back to turbulence. Cloud water was partitioned into liquid and ice with a temperature-dependent scaling factor.

[19] Cumulus clouds were predicted with a modified Arakawa–Schubert algorithm [Ding and Randall, 1998]. In each grid cell, nearly 500 subgrid cumulus clouds could form, each defined by a unique cloud base and top (with up to 23 tropospheric layers, 22 bases and 22 tops were possible). The model accounted for downdrafts and predicted cumulus precipitation, liquid water content, ice content, cumulus cloud fraction, and adjustments to large-scale potential temperature, momentum, and water vapor for each subgrid cloud. The product of each subgrid cloud parameter and subgrid cloud fraction was summed over all subgrid clouds to give bulk values of the parameter for each grid cell.

[20] A free-convective plume model [Lu and Turco, 1994] was applied to each subgrid cumulus cloud to convect gases and aerosols between each subgrid cumulus subcloud layer and cloud top. The resulting vertical gas and aerosol distribution in each subgrid cloud was multiplied by the respective fraction of the subgrid cloud to obtain a weighted-average cloudy-sky vertical gas and aerosol distribution in each grid cell. Cloudy-sky distributions were affected by gas—aerosol—cloud interactions, as described next.

# 2.5. Gas-Aerosol-Cloud Interactions

[21] Size-resolved cloud liquid, ice, graupel, and precipitation in the model interacted with gases and size-resolved aerosol particles, as described here.

- [22] First, summed cloud liquid, ice, and precipitation in each grid cell from the cumulus and stratus schemes, were evaporated/sublimated back to the gas phase so that the vapor could be regrown onto size-resolved ice nuclei (IN) and cloud condensation nuclei (CCN).
- [23] Second, all in-cloud aerosol particle number and component mole concentrations, determined from the convective plume calculation for cumulus clouds and from the cloudy-sky fraction of the grid cell for stratus clouds, were rebinned from the 17-bin aerosol distribution to a 30-bin hydrometeor distribution. A fraction of the particles in each bin were assumed to serve as potential ice nuclei. The fraction was calculated by specifying a probability that a given component in a particle of a given size could serve as an ice nucleus, then weighting the probability by the mole fraction of the component in the bin. The probabilities were estimated from the work of *Pruppacher and Klett* [1997] who suggest that the fraction can be as low as one in a million. They suggest that nonhygroscopic supermicron particles, soil components, a small fraction of OM, certain viruses and bacteria, and a small fraction of sea salt make the best ice nuclei. Soot is not a good ice nucleus, except below  $-15^{\circ}$ C and at high supersaturations. Some pollutants deactivate ice nuclei.
- [24] Third, depositional growth of water vapor to ice was solved simultaneously among the portion of water vapor sublimated from cloud and precipitation total ice and the IN in each of the 30 hydrometeor size bins with a mass-conserving, noniterative and unconditionally stable scheme [*Jacobson*, 1997c]. Whether an IN activated depended on whether it exceeded the critical radius for cloud—ice growth, a prognostic parameter.
- [25] Fourth, growth of water vapor to liquid was solved among the portion of water vapor evaporated from cloud and precipitation total liquid and the CCN in each size bin with the same scheme used for ice growth. The number concentration of CCN in each bin was determined as the total number concentration of aerosol particles minus the number concentration of activated IN. Whether a CCN activated depended on whether its radius exceeded the critical radius for cloud—liquid growth, a prognostic parameter.
- [26] As a result of Steps 1–4, liquid water and ice size distributions were created. Each bin in each distribution contained all the aerosol particle components of the activated CCN or ice nuclei it grew upon. Thus, of the original aerosol particles and their components in each size bin, some were now incorporated within ice crystals, others were incorporated in liquid drops, and the rest were interstitial aerosol particles.
- [27] Fifth, the size-resolved liquid drops and ice crystals were self-coagulated (liquid plus liquid or ice plus ice) to form larger liquid or ice particles and heterocoagulated (liquid plus ice) to form graupel with the scheme described by *Jacobson et al.* [1994] to simulate collision/coalescence. During hydrometeor self-coagulation and heterocoagulation, aerosol particle components incorporated within liquid and ice hydrometeors also coagulated to larger sizes or to the graupel distribution.
- [28] Sixth, the irreversible reactions,  $SO_2(g) + H_2O_2(g) \rightarrow S(VI)(aq)$  and  $SO_2(g) + O_3(g) \rightarrow S(VI)(aq)$  were solved together with gas photochemistry to simulate dissolution and formation of S(VI)(aq) in liquid cloud drops and aerosol

- particles containing liquid water. The rates of these reactions were determined by combining their aqueous rates of reactions with the effective Henry's constant for each species and with the total cloud plus aerosol liquid water content (determined by summing liquid water over all hydrometeor and aerosol size bins). The bulk S(VI)(aq) produced was then distributed over each cloud and aerosol size bin proportionately to the liquid water volume of the bin. The incremental S(VI)(aq) assigned to each bin was added to the S(VI)(aq) already in the bin.
- [29] Seventh, if precipitation occurred from a layer, the largest particles of each hydrometeor type were placed in a separate size distribution and removed from the layer as precipitation along with their associated aerosol particle components, simulating aerosol particle rainout. For each layer, the total mass of liquid water, ice, and graupel precipitated plus the sum of the remaining cloud liquid, ice, and graupel minus the mass of aerosol particles in these distributions exactly equaled the initial mass of bulk cloud plus precipitation liquid and ice from the cumulus and stratus parameterizations.
- [30] Eighth, size-resolved interstitial aerosol particles and their components were heterocoagulated (again conserving volume) with each of the three size-resolved cloud hydrometeor distributions in each layer and with each of the three size-resolved precipitation hydrometeor distributions passing through each layer. The latter allowed treatment of aerosol particle washout (rainout was treated in Step 7). The precipitation passing through each layer was the sum of precipitation from all cloud layers above.
- [31] Ninth, the precipitation passing through each layer was summed over all hydrometeor sizes to obtain the bulk precipitation passing through the layer. Equilibrium equations between each gas in the model and the bulk rainwater (assuming an effective Henry's constant for each gas) were solved to estimate the mole fraction of the gas from each cell partitioned into rainwater, simulating gas washout. If rainwater passing through a cell was already saturated with the gas, no additional gas could enter the rainwater.
- [32] Finally, at the end of each time step the nonprecipitated cloud liquid, ice, and graupel distributions were evaporated, releasing the aerosol particle cores. The evaporated aerosol particle distribution differed from the original in-cloud distribution due to in-cloud aerosol particle growth, coagulation, chemistry, evaporation, and removal. The evaporated distributions were then weighed by cloud fraction and added to clear-sky distributions, weighted by clear-sky fraction. The liquid water content of the resulting aerosol particles was determined from an equilibrium calculation (section 2.2).

# 2.6. Radiative Transfer

[33] Radiative transfer was solved to determine heating rates (for temperature calculations) and actinic fluxes. Irradiances for heating rates were determined for 409 wave intervals (86, 67, and 256 in the regions <0.8, 0.8–4.5, and 4.5–1000  $\mu$ m, respectively). Actinic fluxes were determined for 86 intervals <0.8  $\mu$ m. Both were solved with the technique of *Toon et al.* [1989], which has an error <10% under typical scattering and absorbing conditions. In each column of the model, two calculations were performed: one with clouds, aerosols, and gases and the other

with aerosols and gases, but no clouds. Heating rates and actinic fluxes from the two calculations were then weighted by cloudy-sky and clear-sky fractions. Optical properties used for radiative transfer are described as follows.

[34] The model accounted for Rayleigh scattering and gas absorption. UV and/or visible light gas absorbers included O<sub>2</sub>, O<sub>3</sub>, CO<sub>2</sub>, H<sub>2</sub>O, NO<sub>2</sub>, NO<sub>3</sub>, N<sub>2</sub>O, N<sub>2</sub>O<sub>5</sub>, HONO, HNO<sub>3</sub>, HO<sub>2</sub>NO<sub>2</sub>, H<sub>2</sub>O<sub>2</sub>, HCHO, CH<sub>3</sub>CHO, and CH<sub>3</sub>OOH. Solar-IR absorbers included H<sub>2</sub>O, CO<sub>2</sub>, O<sub>3</sub>, and O<sub>2</sub>. Thermal-IR absorbers included H<sub>2</sub>O, CO<sub>2</sub>, O<sub>3</sub>, CH<sub>4</sub>, and N<sub>2</sub>O. Thermal-IR absorption coefficients were from the work of *Mlawer et al.* [1997].

[35] Aerosol particle optical properties were calculated assuming that BC (if present) comprised a particle's core and all other material coated the core. BC density was 1.5 g/ cm<sup>3</sup>, and its refractive indices were taken from the work of Krekov [1993]. The ratio of BC's midvisible imaginary refractive index to its density was less than that of crystalline graphite, suggesting that results here may underpredict BC radiative effects. Shell refractive indices were obtained by calculating a solution and nonsolution refractive index and volume averaging the two. Solution real refractive indices were calculated from partial-molar refraction theory [e.g., Stelson, 1990; Tang, 1997]. Solution imaginary refractive indices were calculated analogously. Refractive indices for OM (a nonsolution component) were obtained from the work of Krekov [1993], except that UV refractive indices were weighted to account for selective UV absorption by nitrated aromatic, polycyclic aromatic, and other UVabsorbing organic compounds, as described by Jacobson [2001b]. Core and shell refractive indices were used in a core-shell Mie theory calculation [Toon and Ackerman, 1981] to determine optical properties for each particle composition and size and for each wavelength.

[36] Cloud liquid, ice, and graupel optical properties for each hydrometeor size and radiation wavelength were also determined from Mie calculations.

# 2.7. Subgrid Temperatures and Fluxes and Ocean Treatment

[37] The model treated ground temperatures over heterogeneous surfaces. Each grid cell was divided into up to 15 surface classes (12 soil classes and roads, roofs, and water). The model also treated vegetation over soil, snow over bare soil, snow over vegetation over soil, sea ice over water, and snow over sea ice over water. For surfaces except sea ice and water, temperatures and liquid water were found with a timedependent 10-layer subsurface module. For sea ice, a timedependent slab calculation was performed. For ocean water, a 2-D potential-enstrophy conserving shallow-water equation mixed-layer module, forced by wind stress (G. Ketefian, Ph.D. thesis, Stanford University, 2002, in preparation, based on the shallow-water scheme of Arakawa and Lamb [1981]) was used to predict ocean velocities and transport energy horizontally. The average mixing depth was 60 m although the actual depth at each location was a prognostic variable. For lake water, a fixed 60 m mixing depth was assumed. Water temperatures were also affected by sensible, latent, and radiative fluxes between water and air.

[38] At the end of a subsurface-module time step (12 s), surface temperatures, sensible heat fluxes, latent heat fluxes, water vapor fluxes, friction velocities, surface albedos,

emissivities, and other parameters were weighted by the fractional surface type in the cell to give an effective parameter value for the cell as a whole. The model treatment of surfaces is discussed in detail by *Jacobson* [2001c], except that global soil moisture was initialized here with monthly gridded values from the work of *Nijssen et al.* [2001] and the global fractional vegetation cover was obtained here from the work of *Zeng et al.* [2000]. The surface albedos for snow and sea ice used here were wavelength dependent [*Grenfell et al.*, 1994; *Allison et al.*, 1993] and a function of snow and/or sea ice thickness.

### 3. Effects of Particles on Climate

[39] The model treated 12 identifiable effects of aerosol particles on climate. Two of these effects (the "indirect effect" and the "semidirect effect") have previously been specified in the literature. Two additional effects (the "particle effect through surface albedo" and the "particle effect through large-scale meteorology") have obvious counterparts with respect to greenhouse gases and have been accounted for in numerous previous studies. The remaining eight effects have either not been described or described only partially in the literature. Some have been accounted for in simulations.

#### 3.1. The "Self-Feedback Effect"

[40] When new particles are emitted, they affect air temperature, relative humidity, and surface area available for gases to condense upon, all of which affect the composition, liquid water content, size, and optical properties of both the new and existing particles. This effect is referred to as the "self-feedback effect." For example, when BC warms the air, it decreases the relative humidity, decreasing the liquid water content and reflectivity of particles containing sulfate and nitrate, warming the air further. At the same time, when BC is emitted in one location, it provides additional surface area for sulfuric acid to condense upon, increasing the formation of sulfate upwind and decreasing it downwind. An increase in sulfate increases its liquid water content as well. Changes in aerosol particle liquid-water also change the aqueous oxidation rate of SO<sub>2</sub> to sulfate and the dissolution rate of NH<sub>3</sub>, HNO<sub>3</sub>, and HCl into particles, further changing particle size and reflectivity. Treatment of the "self-feedback effect", which requires the treatment of nucleation, electrolyte hydration, dissolutional growth, condensational growth, aqueous chemistry, and the effect of aerosol particles on temperatures and the relative humidity through the thermodynamic energy equation has been accounted for in model simulations [*Jacobson*, 1994, 1997a, 1998a].

### 3.2. The "Photochemistry Effect"

[41] Aerosol particles alter photolysis coefficients of gases, affecting their concentrations and those of other gases (through kinetic chemistry) [e.g., Jacobson, 1994, 1997b, 1998b; Dickerson et al., 1997]. A change in gas abundance changes solar and thermal-IR heating rates in a process called the "photochemistry effect". The effect of aerosols on heating rates through their modification of photolysis has been accounted for in 3-D simulations [Jacobson, 1994, 1997b, 1998b]. To account for the "photochemistry effect", changes in size-resolved particles were

allowed to affect photolysis and solar plus thermal-IR heating rates of all UV- and solar-absorbing gases and the heating rates of their by-products.

## 3.3. The "Smudge-Pot Effect"

[42] During day and night all aerosol particles trap the Earth's thermal-IR radiation, warming the air [Bergstrom and Viskanta, 1973; Zdundowski et al., 1976]. This warming is well known to citrus growers who, at night, used to burn crude oil in smudge pots to fill the air with smoke and trap thermal-IR radiation, preventing crops from freezing. The warming of air relative to a surface below increases the stability of air, reducing vertical fluxes of horizontal momentum, slowing surface winds (and increasing them aloft), reducing the wind speed-dependent emission rates of sea-spray, soil-dust, road-dust, pollens, spores and some gas-phase particle precursors. The reduction in these particles affects daytime solar and daytime/nighttime thermal-IR radiation transfer. Changes in stability and winds also affect energy and pollutant transport. The effect of thermal-IR absorption by particles on the emission of other particles and gases and on local energy and pollutant transport is referred to as the "smudge-pot effect". In the model, aerosol particle scattering and absorption affect solar and thermal-IR heating rates, which change temperatures and stability, which feed back to winds, emissions of winddriven aerosol particles and precursor gases, and pollutant/ energy transport, allowing for the treatment of this effect.

#### 3.4. The "Daytime Stability Effect"

[43] If airborne particles absorb solar radiation, the air warms. Whether the particles absorb or only scatter, they prevent solar radiation from reaching the surface, cooling the surface and increasing the air's stability [Bergstrom and Viskanta, 1973; Venkatram and Viskanta, 1977; Ackerman, 1977]. Like with the "smudge-pot effect", enhanced daytime stability slows surface winds, reducing emissions of wind-driven particles and gases and affecting local pollutant and energy transport. At the same time, heating at the top of the boundary layer due to aerosol absorption destabilizes the air above the boundary layer [e.g., Jacobson, 1998b, Figure 8]. This destabilization has little effect on emissions, since the air below is more stable, but slightly increases venting of particles at the top of the boundary layer to the free troposphere. The effect of solar absorption and scattering by particles on emissions of other particles and gases and on local energy and pollutant transport is referred to as the "daytime stability effect". The same processes in the model that allow treatment of the "smudge-pot effect" allow treatment of this effect.

#### 3.5. "Indirect Effects"

[44] An increase in the number of particles increases the number and decreases the size of cloud drops ("first indirect effect") [Twomey, 1977], reducing rates of drizzle, thereby increasing liquid water content and fractional cloudiness of low-level clouds ("second indirect effect") [Albrecht, 1989]. These "indirect effects" increase scattering and decrease the transmission of radiation to the Earth's surface. In the model, clouds formed on top of current aerosol particle size distributions. Size-resolved cloud drops and ice crystals grew by condensation, deposition, and coagu-

lation, and the largest resulting drops, crystals, and graupel were removed by precipitation, allowing for substantial treatment of the "indirect effects."

# 3.6. The "The Effect on BC Absorption of the First Indirect Effect"

[45] When the "first indirect effect" occurs, cloud scattering is enhanced, increasing the absorption of all solar radiation by BC and solar-IR radiation by water vapor and CO<sub>2</sub> within and above the cloud, warming the air there. Previous studies have shown that clouds enhance the direct forcing of soot lying within and above them [e.g., *Haywood et al.*, 1997; *Jacobson*, 2001b], but studies to date have not linked changes in cloud scattering due to the first indirect effect to enhanced absorption by BC or greenhouse gases.

#### 3.7. The "Semidirect Effect"

[46] Solar absorption by a low cloud increases stability below the cloud, reducing vertical mixing of moisture to the cloud base, thinning the cloud [Nicholls, 1984]. Decreases in relative humidity correlate with decreases in low-cloud cover [Bretherton et al., 1995; Klein, 1997]. Similarly, absorbing particles warm the air, decreasing its relative humidity and increasing its stability, reducing low-cloud cover [Hansen et al., 1997; Ackerman et al., 2000] Reduced cloud cover increases sunlight reaching the surface, warming the surface in a process called the "semidirect effect" [Hansen et al., 1997]. In the model, in-cloud BC absorption of solar radiation allowed for this effect.

### 3.8. The "BC-Low-Cloud Positive Feedback Loop"

[47] When BC reduces low-cloud cover by increasing stability and decreasing relative humidity, enhanced sunlight through the air is absorbed by BC (and by water vapor and  $\mathrm{CO}_2$  in the solar-IR), further heating the air and reducing cloud cover in a positive feedback loop, identified here as the "BC-low-cloud positive feedback loop". Whereas  $\mathrm{CO}_2$  also warms the air by absorbing thermal-IR and solar-IR radiation, reducing low cloud cover and enhancing surface solar radiation in some cases, it absorbs less sunlight than BC, so  $\mathrm{CO}_2$  partake less in this feedback loop. The model allowed treatment of this effect, since solar absorption by BC,  $\mathrm{CO}_2$ , and  $\mathrm{H}_2\mathrm{O}$  was affected by cloud optics.

## 3.9. The "Rainout Effect"

[48] When BC increases stability of and vertical moisture transport in the boundary layer, it may decrease cumulus convection, much of which starts in the boundary layer, reducing cumulus precipitation. The addition of BC also triggers the second "indirect effect," which reduces precipitation. Reducing precipitation reduces rainout of soluble gases and all aerosol particles, increasing their concentrations in the air, increasing warming in some cases and cooling in others.

### 3.10. The "BC-Water Vapor Positive Feedback"

[49] When BC is present, it warms the air relative to the surface. The warmer air decreases the relative humidity, evaporating cloud water to vapor, a greenhouse gas, warming the air further. The resulting vertical temperature gradient, in the presence of mechanical or thermal turbulence, increases the downward sensible heat flux, warming the surface (although less than the air is warmed). The increase

in surface temperature relative to its initial temperature increases the evaporation rate of ocean and soil water. Since the air is now warmer, the saturation vapor pressure of water is now higher, allowing much of the additional water vapor to accumulate. The additional water vapor absorbs solar-IR and thermal-IR, warming the air further. (This feedback also applies to  $\mathrm{CO}_2$  for the most part).

### 3.11. The "Particle Effect Through Surface Albedo"

[50] During the day, airborne BC reduces sunlight to the ground, cooling it, increasing the lifetime of existing snow. Conversely, because BC warms the air, snow falling through the warmer air is more likely to melt. At night, airborne BC also enhances downward thermal-IR, increasing nighttime melting and sublimation of snow on the ground. Because the albedo of new snow exceeds that of sea ice, which exceeds those of soil or water, the melting of snow or sea ice increases sunlight to the surface. The effect of aerosol particles on temperatures through their change in snow and sea ice cover is referred as the "particle effect through surface albedo", analogous to its well-known greenhouse gas counterpart, the "snow-albedo effect". Any model that treats the climate response of aerosols treats this effect to some degree. A related effect is the effect of soot deposition to snow and sea ice on their albedos [e.g., Warren, 1984; Vogelmann et al., 1988; Warren and Clarke, 1990; Grenfell et al., 1994; Light et al., 1998; Hansen and Sato, 2001]. This effect was not treated here. It may be more important where pollutant concentrations are high [e.g., Light et al., 1998] than over the Antarctic [e.g., Warren and Clarke, 1990; Grenfell et al., 1994].

# 3.12. The "Particle Effect Through Large-Scale Meteorology"

[51] Aerosol particles affect local temperatures, which affect local air pressures, winds, relative humidities, and clouds. Changes in local meteorology slightly shift the locations and magnitudes of semipermanent and thermal pressure systems and jet streams. The effect of local particles on large-scale temperatures is referred to as the "particle effect through large-scale meteorology" and has a well-known counterpart with respect to greenhouse gases. Any model that treats the climate response of aerosols treats this effect.

# **3.13.** Difficulty in Isolating the Importance of the Effects

- [52] Whereas, it would be ideal to quantify the importance of each effect described, this is not possible nor has it been done for any effect to date. Quantifying any effect of aerosol particles requires shutting off one or more physical or chemical processes; yet, each chemical and physical aerosol processes affects several effects simultaneously, not just one, so it is possible only to quantify correctly the net effect of all feedbacks on climate simultaneously, but not an individual feedback.
- [53] For example, previous studies have tried to quantify the "indirect effects". In the case of the "first indirect effect" they have calculated the difference in forcing from two simulations, one with present-day CCN and another with preindustrial CCN [e.g., *IPCC*, 2001]. Such a comparison does not (nor can any comparison) correctly quantify the

"first indirect effect", since in the real atmosphere, changing the number of CCN immediately triggers the "photochemistry effect", the "effect on absorption of the first indirect effect", the "smudge-pot effect", the "daytime stability effect", the "semidirect effect", the "BC-low-cloud positive feedback loop", etc., all of which feed back to temperatures and to the number of CCN. In all cases, either these models did not account for some or all of these effects, in which case their climate responses and estimated forcings were incorrect, or they really quantified the sum of multiple effects. More specifically, if a study wants to isolate the "indirect effects" due to BC correctly, it must account for all effects that affect BC (and all other aerosol) concentrations and coatings, since these parameters affect the "indirect effects" themselves. Yet, a study that includes these other effects cannot separate out their climate responses from those of the "indirect effects". Although they cannot be correctly quantified, effects can be used to explain some characteristics of climate response, as described in section 6.

### 4. Simulations

- [54] Three sets of simulations were run: with and without (1) current levels of anthropogenic CO<sub>2</sub>, (2) current levels of anthropogenic CH<sub>4</sub>, and (3) emitted f.f. BC + OM. f.f. BC represents about half of global anthropogenic BC. The other half is from biomass burning. Biomass-burning emissions were included here, but this paper examines the climate response of reducing only f.f. BC + OM emissions, not biomass-burning emissions.
- [55] The with/without anthropogenic CO<sub>2</sub> and CH<sub>4</sub> simulations were "equilibrium climate sensitivity" simulations [IPCC, 2001] comparing current and preindustrial CO<sub>2</sub> and CH<sub>4</sub> loadings, respectively. The equilibrium simulations were each run for 6 years, more than the estimated equilibrium response time of temperature to a step change in forcing in a model with a mixed-layer ocean (section 2.7) [e.g., Hoffert et al., 1980]. A comparison of results from the equilibrium CO<sub>2</sub> simulations with radiosonde measurements and another model study is discussed in section 6. In addition, a 6 year CO<sub>2</sub> doubling equilibrium sensitivity was also run, and it caused a warming of 3.2 K, comparable with IPCC [2001] CO<sub>2</sub> doubling sensitivities of 1.5–4.5 K.
- [56] The historic CO<sub>2</sub> and CH<sub>4</sub> equilibrium simulations were run to estimate the effect of eliminating all anthropogenic emissions of CO2 and CH4. Even if all anthropogenic CO<sub>2</sub> emissions were eliminated today, full climate response would not occur for 1000 years, since the e-folding lifetime of CO<sub>2</sub> is 50–200 years [IPCC, 2001], and several lifetimes are required for a perturbation to be cleared out of the atmosphere. Figure 1 scales the equilibrium results with lifetime to show how eliminating CO<sub>2</sub> or CH<sub>4</sub> emissions today might effect global near-surface temperatures over the next 100 years. The use of a mixed-layer ocean model (section 2.7) assumes that the deep ocean has little effect on temperatures over the 100 year period. Although this is a simplifying assumption, Figure 9.1 of IPCC [2001] suggests that the deep ocean may have little net effect on atmospheric temperatures due to perturbations to CO2 over 80 years, and between years 80 and 100, the deep ocean dampens temperature changes due to CO<sub>2</sub> by about 7%. Such a dampening would appear to shift all curves in

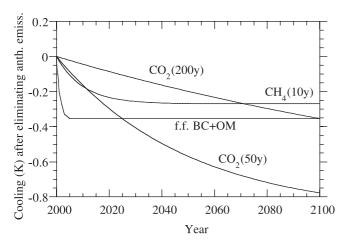


Figure 1. Estimated changes in temperature due to eliminating all anthropogenic emissions of each CO<sub>2</sub>, CH<sub>4</sub>, and f.f. BC + OM during the next 100 years. The curve for CO<sub>2</sub> and CH<sub>4</sub> were obtained by first running "equilibrium climate sensitivity" simulations with and without current levels of anthropogenic mixing ratios of each component for 6 years. The differences in temperatures between the with and the without cases  $(-0.9 \text{ K for CO}_2,$ -0.27 K for CH<sub>4</sub>) represent the cooling that results after eliminating the component from the atmosphere, which would occur only after many e-folding lifetimes of the component. The differences in temperatures were exponentially decayed from the initial time, assuming initial efolding lifetimes of 50 and 200 years for CO<sub>2</sub> [curves CO<sub>2</sub> (50 years) and CO<sub>2</sub> (200 years), respectively] and 10 years for CH<sub>4</sub> [curve CH<sub>4</sub> (10 years)]. In the case of f.f. BC + OM, simulations were run with/without f.f. BC + OM emissions (5.1 Tg/yr BC + 10.1 Tg/yr OM) for 5 years. The curve after 5 years was held constant.

Figure 1 up proportionately after 80 years, thereby strengthening the conclusions here, since the largest absolute changes would occur for CO<sub>2</sub>. Similarly, if the deep ocean affected atmospheric temperatures more prior to 80 years, the effect would apply proportionally to all warming components, possibly strengthening the conclusions here with respect to f.f. BC + OM.

[57] Time-dependent simulations with and without f.f. BC + OM emissions were run because it is difficult to control BC without controlling OM too. For the baseline simulation, emissions of all gas and aerosol species described in sections 2.1 and 2.2, were included, and all processes described in section 2 were solved. Due to the computational burden of the simulations (approximately 35 days of computer time per simulation year), only 5 years of simulation were performed and results were held constant thereafter. For the sensitivity simulation, the only change made was that f.f. BC + OM emissions were removed. Major f.f. sources of BC include hard coal, brown coal, diesel fuel, and jet fuel. The ratios of emitted OM/BC from these fuels are about 1.9:1, 3.9:1, 1:1, and 0.6:1, respectively, giving a weighted ratio of about 2:1 [Cooke et al., 1999] [assuming an OM/OC ratio of 1.3]. The 2:1 ratio was used here when removing OM and BC from f.f. Sulfate now comprises <1% of diesel emissions [Shauer et al., 1999].

[58] Figure 1 shows that, after 5 years of the transient simulation, removing f.f. BC + OM cooled climate by about 0.35 K. The cooling exceeded that from removing CH<sub>4</sub> emissions for all 100 years and CO<sub>2</sub> emissions for 25–100 years (depending on whether CO<sub>2</sub>'s lifetime was 50 or 200 years). The figure further implies that any emission reduction of f.f. BC + OM would cool climate more than would any emission reduction of CO<sub>2</sub> or CH<sub>4</sub> for a specific period. The period approaches zero in the extreme case of a minute reduction in f.f. BC + OM and a complete reduction in anthropogenic CO<sub>2</sub>.

[59] The sum of warming due to anthropogenic  $CO_2 + CH_4 + f.f.$  BC + OM was +1.5 K. Other greenhouse gases, if scalable to  $CO_2$ , cause +0.4 K warming, bringing the total warming to +1.9 K. Observed warming from 1850 to 2000 is +0.75  $\pm$  0.2 K [Jones et al., 1999]. It is hypothesized that the 1.2 K difference is due primarily to cooling by anthropogenic particles other than f.f. BC + OM, such as sulfate, nitrate, secondary organics, fly ash, soil dust, and reflective anthropogenic biomass-burning particles. Natural climate variations could also be responsible for some of the difference, but the net historic effect between 1860 and 2000 may be small [Stott et al., 2000].

[60] To test the hypothesis that non-BC aerosols cause cooling, time-dependent simulations (accounting for all effects discussed) in which all f.f. SO<sub>2</sub> emissions were and were not included were run. The short-term cooling due to f.f. SO<sub>2</sub> alone was -0.7 K. Further simulations are necessary to elucidate climate responses of other aerosol particle components, but given the magnitude of the atmospheric burdens of these other components, it appears plausible that they could be responsible for the additional cooling.

[61] In sum, the observed 1850–2000 warming can feasibly be reconciled by considering greenhouse gases, f.f. BC + OM, and other particulate, implying that non-BC + OM particulates mask much actual global warming to date. The results here also imply that eliminating all f.f. BC + OM with no other change might reduce >40% (0.35 K/0.75 K) of net global warming to date and >15% (0.35 K/1.9 K) of total global warming before cooling is subtracted out. Based on sensitivity tests, the estimated ranges of these values are 20–45% and 8–18%, respectively. Reducing CO<sub>2</sub> emissions by a third would have the same effect, but over 50–200 years.

[62] The long-term climate responses of reducing emissions calculated here (-0.35 K for f.f. BC + OM, -0.9 K for CO<sub>2</sub>, -0.27 K for CH<sub>4</sub>, and +0.7 K for SO<sub>2</sub>) are within uncertainty ranges of those calculated using a different approach by *Hansen and Sato* [2001]. Their values were -0.38 K for f.f. BC, -1.05 K for CO<sub>2</sub>, -0.53 K for CH<sub>4</sub>, and +0.75 K for SO<sub>2</sub>. Their approach was to estimate a total (direct plus other) forcing for each component and multiply the estimated total forcing by a climate response function, 0.75 K per W/m<sup>2</sup> total forcing.

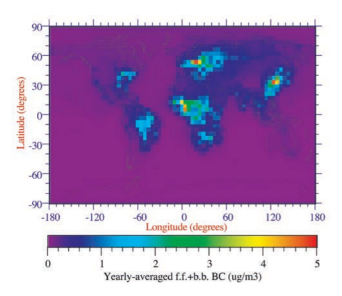
[63] Another type of climate response function is the climate (K) response per unit direct forcing rather than total forcing. For this study, these functions were 1.4 (0.35 K/ $0.25 \text{ W/m}^2$ ), 0.6 (0.9 K/ $1.6 \text{ W/m}^2$ ), 0.6 (0.27 K/ $0.47 \text{ W/m}^2$ ), and 2 ( $-0.7 \text{ K/}-0.35 \text{ W/m}^2$ ) for f.f. BC + OM, CO<sub>2</sub>, CH<sub>4</sub>, and SO<sub>2</sub>, respectively. The f.f. BC + OM and SO<sub>2</sub> ratios were about 2 and 3 times higher, respectively, than were the CO<sub>2</sub> and CH<sub>4</sub> ratios, because particles have several effects that major greenhouse gases do not (e.g., "indirect effect,"

"self-feedback effect", "semidirect effect", "BC-low-cloud positive feedback loop", photochemistry effect", and "rainout effect"). SO2 had a greater climate response (in the opposite direction) than did BC + OM, partly because the 'indirect effect" cooled climate in both cases, offsetting some warming in the case of BC + OM and increasing cooling in the case of SO<sub>2</sub>. Also, because SO<sub>2</sub> is a gas and spreads further than does BC + OM, the resulting sulfate spread further and had a greater hemispheric "indirect effect" than did BC + OM. Results here indicate that sulfate increased cloud optical depths more than did BC + OM. The 3:1 ratio of sulfate to CO<sub>2</sub> climate response divided by direct forcing is in the range of IPCC [2001], who assumes that temperatures are proportional to total forcing and suggest that the ratio of the aerosol indirect "forcing" upper limit plus sulfate direct forcing mean to the sulfate direct forcing mean is 6:1, compared with 1:1 for CO<sub>2</sub>.

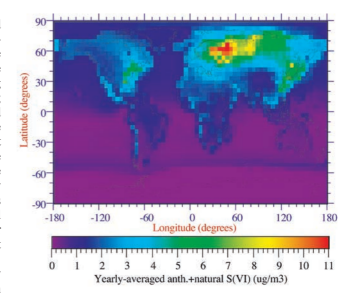
[64] Results here suggest that BC warmed the air by 360,000-840,000 times more effectively per unit mass than did CO<sub>2</sub>, which illustrates why a small mass of BC can have such a large climate effect. In the simulations, the global loading of f.f. BC + OM (where the OM in this term is primary) varied during the year from 0.25 to 0.6 Tg (one-third was f.f. BC and two thirds f.f. OM). The atmospheric loading of anthropogenic CO<sub>2</sub> is approximately 182,000 Tg. Dividing the long-term warming due to f.f. BC + OM (0.35 K) and anthropogenic CO<sub>2</sub> (0.9 K) by their respective loadings and dividing the result for f.f. BC + OM by that for CO<sub>2</sub> gives the warming per unit mass of f.f. BC + OM versus that of CO<sub>2</sub> as 120,000-280,000:1. Multiplying by three (the mass ratio of BC + OM to BC) gives the warming per unit mass of f.f. BC versus that of CO<sub>2</sub> as 360,000-840,000:1.

# 5. Uncertainties and Comparisons With Observations

[65] Uncertainties in the model include uncertainties in continuous input data (e.g., emission data), model data (e.g.,



**Figure 2.** Fifth-year average global distribution of modeled near-surface biomass-burning plus f.f. BC  $(\mu g/m^3)$ , obtained from the baseline simulation.



**Figure 3.** Fifth-year average global distribution of modeled near-surface anthropogenic plus natural S(VI)(aq) (μg/m³), obtained from the baseline simulation.

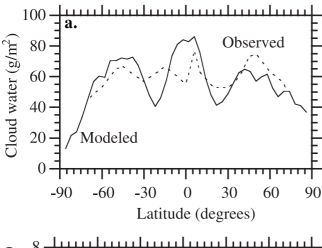
chemical rate coefficient, absorption cross section, aerosol thermodynamic, refractive index, land use, and soil data), model processes, model resolution, and model numerics.

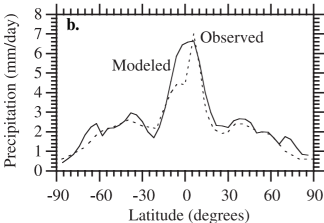
- [66] A process that was ignored (in part due to the short simulation period) was long-term natural climate variation. This process would affect predictions over the long term, but should not affect the conclusions here, since, natural variations are expected to affect the climate relatively equally when an anthropogenic component in present and absent.
- [67] Another source of uncertainty is random model variability. For example, would the results change if initial conditions changed or a model process was treated differently. To test this, 11 additional simulations with and without f.f. BC + OM were performed, each with a different initial condition, emissions, or treatment of one or more model processes. The simulations resulted in net globally averaged temperature changes due to f.f. BC + OM of +0.15 to +0.5K (compared with the +0.35 K from the baseline case), indicating some variability in predictions of the climate response of f.f. BC + O but in all cases supporting the conclusions found here.
- [68] Figure 2 shows modeled near-surface BC concentrations from the baseline simulation. BC sources included biomass burning and f.f. Table 1 compares modeled versus measured near-surface BC. The table indicates that BC concentrations were, on average, slightly higher than observations over marine regions but much lower than observations over many urban areas, where BC concentrations were highest, suggesting that BC warming may be higher than estimated here.
- [69] Figure 3 shows modeled near-surface anthropogenic plus natural S(VI) concentrations from the baseline case. Table 2 compares modeled with measured S(VI). S(VI) predictions compared well in many urban and rural areas, but they were low in some of the most polluted areas (e.g., Beijing, Nylsvley, Atlanta, and Po Valley). Sulfate was somewhat overpredicted in a few remote areas and underpredicted in others.

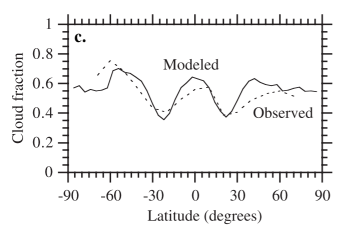
	ions	
	trat	
	cen	
	Conc	
١	o	
	tate	
	Ħ	
	n	
	ace	
	nrta	
	Ŋ	
	Near-Surtac	
	Z	
	J	
	rved	
	ser	
	Obser	
(	ゔ	
	sns	
	ers	
	>	
	leled	
	g	
	ğ	
	$\geq$	
	ot Mo	
	mparison	
	18	
	ä	
	ď	
	5	
	$\mathcal{L}$	
	ri	
	Table 7	
	<u>ड</u>	
	ಡ	

January—February  April—October March—June Annual December—February Annual April—May July—August Annual	Chapter I address I address	Lotitudo	Jensie and I	Modeled milfets	Macanimod militato	D	Dots actions
Coem         31–37%         66 – 76°W         3.00         Marine         1005–540         January-February           8 – 15 N         8 – 15 N         33–46°W         5.60         5.60         1.60–2400         January-February           8 – 15 N         8 – 15 N         33–46°W         5.60         8.00         1.60–2400         January-February           9 – 5 N         17 – 26°W         2.60         8.00         1.60         3.00         January-February           1 S – 48°N         18 – 6°E         8.00         1.60         8.00         January-February           1 S – 48°N         1 S – 6°E         8.00         1.17         January-February           1 S – 48°N         1 S – 6°E         8.00         1.17         January-February           1 S – 48°N         1 S – 6°E         8.00         January-February           1 S – 48°N         1 S – 6°E         8.00         January-February           Abrual Land, Aniactica         1 S – 6°E         1.00         January-February           Abrual Land, Aniactica         1 S – 6°E         1.00         January-February           Abrual Land, Aniactica         1 S – 6°E         1.00         January-February           Abrual Land, Aniactica         1 S – 6°E <td< th=""><th>Station</th><th>Lautude</th><th>Longinae</th><th><math>(ng/m^3)</math></th><th><math>(ng/m^3)</math></th><th>reitou</th><th>Data reference</th></td<>	Station	Lautude	Longinae	$(ng/m^3)$	$(ng/m^3)$	reitou	Data reference
Cocum         13-17N         66-76°W         2400         1000-1800         January-February           15-137N         64-66°W         550         100-1800         January-February           15-137N         63-48°W         550         100-1800         January-February           3-8°N         33-48°N         15-24°W         410         880-2100         Amual           3-8°S-3'N         17-24°N         150         685-1720         Amual           66'S-18         15-24°N         150         880-2100         Amual           Antactica         68'S-18         150         880-2400         Amual           Antactica         68'S-18         152         190         Amual           Antactica         78'S-18         1152-2°W         30         208         Amual           Antactica         78'S-18         1152-2°W         30         208         Amual           Antactica         78'S-18         1152-2°W         30         20         Amual           Antactica         68'S-18         110         30         Amual         Amual           Antactica         1150         110         110         Amual         Amual           Antactica         111 <td></td> <td></td> <td></td> <td></td> <td></td> <td></td> <td></td>							
15-21'N	Atlantic Ocean	$31-37^{\circ}N$	$M_{\odot}92-99$	2400	1060 - 540	January-February	<i>Quinn et al.</i> [2001]
17.2   17.2		$15-31^{\circ}N$	$43-66^{\circ}$ W	550	500 - 1080		
5°S - 3°N         2°S - 3°N         2°S - 3°N         410         800 - 2400           A. Antarctical         5°S - 3°N         1 - 26°N         410         800 - 2400           A. Antarctical         2°S - 3°N         1 - 26°N         410         800 - 2400           file         31 - 24°N         1 - 26°N         85         65 - 1720         Armual           file         27 - 48°N         1   15°2°         85         60 - 18°0         Armual           mand         Actic         878°         1   15°3°         Armual         Armual           mand         Actic         1   15°3°         45°         90         Armual           mand         Actic         1   15°3°         1   15°3°         Armual           Abatractical         1   15°3°         1   15°3°         Armual           Abatractic         1   10°7°         1   10°7°         Armual           Abatractic <t< td=""><td></td><td><math>8-15^{\circ}{ m N}</math></td><td><math>33-43^{\circ}</math>W</td><td>560</td><td>1760 - 2490</td><td></td><td></td></t<>		$8-15^{\circ}{ m N}$	$33-43^{\circ}$ W	560	1760 - 2490		
And Antercical Controls         55-5-78 (PL-17°W)         250         880-2100           And arterical Controls         53-24°S         67-17°W         550         880-2100           fife         86°S         66-1720         Ammal           fife         87         96-1720         Ammal           sund Acrie         78°S4         17.3°F         120         500-4000         April-June           Amarctica         78°S4         118°5F         120         500-4000         April-June           Amarctica         78°S4         118°3F         30         51         April-June           Amarctica         78°S4         143°F         2490         150         December           Amarctica         78°S4         119°T         2490         150         December           in         118°T         60°T         250         155         December           in         118°T         55°         250         155         December           in         3°N         60°N         38°         40°         37°         April-May           in         3°N         69°W         88°         25°         40°         April-May           in         3°N         <		$3-8^{\circ}N$	$26 - 33^{\circ} \text{W}$	410	890-2400		
n, Antarctical         52-5°         9° E-17°         150         650-1570           n, Antarctical         68°S         16° E         150         650-1570           n, Antarctical         68°S         11° SE         150         650-1570           n, Antarctical         68°S         11° SE         100         500-1570         Antanual           and, Actic         77° 48°         11° SE         10°         500-170         April-October           and, Actic         78°S         11° SE         10°         500-170         April-October           and, Actic         78°S         11° SE         10°         500-00         April-October           Antancetical         18°S         11° SE         10°         50°         April-October           Antancetical         18°S         11° SE         10°         April-October         April-October           Antancetical         41°         41°         42°         42°         April-October           Antancetical         41°         42°         42°         April-October         April-October           Antancetical         41°         42°         42°         42°         April-October           Antancetical         41°         42°		$5^{\circ}\text{S}-3^{\circ}\text{N}$	$17-26^{\circ}{ m W}$	250	880-2100		
3.2-4.5         18-6°E         85.0         965-1720         Annual           diff         68.5         66.5         17.0         500-4000         April-October           life         27-48°N         175°E         190         500-4000         April-October           hife         27-48°N         1197°E         190         April-October         April-October           Antarctica         66°S         143°S         144°E         29.0         1960         December - February           Antarctica         66°S         144°E         240         1960         December - February           Antarctica         37°S         119°T         240         1960         December - February           Antarctica         43°S         119°T         240         1960         December - February           Antarctica         43°S         119°T         240         1960         December - February           Antarctica         43°S         119°T         240         1960         December - February           Antarctica         113°N         175°N         26°D         250         140°D         April-May           Antarctica         47°N         44°N         14°C         25°D         44°N <td< td=""><td></td><td><math>25-5^{\circ}S</math></td><td><math>6^{\circ}\mathrm{E}{-}17^{\circ}\mathrm{W}</math></td><td>150</td><td>630 - 1570</td><td></td><td></td></td<>		$25-5^{\circ}S$	$6^{\circ}\mathrm{E}{-}17^{\circ}\mathrm{W}$	150	630 - 1570		
tip         50°-40°         68°-E         30         1112         Annual           itip         27-48°N         178°E         30         500-4000         Annual           sund, Antarctic         76°S         64°W         45         97         Annual           Antarctic         76°S         11°33E         30         228         December - February           Antarctica         77°S S         11°28W         240         1960         Annual           Antarctica         43°S I         14°S S         197         Annual           Antarctica         43°S I         11°SW         240         1960         Annual           Antarctica         43°S I         14°S I         14°S I         Annual           Inbuic         41°N I         55°W I         25°0         Annual           Infuir         47°N I         56°W I         38°0         25°0         Annual           Infuir         47°N I         17°3         48°0         48°3         Annual           Infuir         47°N I         17°3         48°3         Annual           Infuir         47°N I         17°3         48°3         Annual           Infuir         47°N I         17°		$33-24^{\circ}S$	$18-6^{\circ}\mathrm{E}$	850	965-1720		
tild         77–48° N         113% E         100         500–4000         April—Cotober Amunal Amunchan (5°% Amunal Amunal Amunchan (5°% Amunal	Mawson, Antarctica	S.89	63 °E	30	112	Annual	Savoie et al. [1993]
Antacctican distance         678° 54" (1°53°F	NE Pacific	$27-48^{\circ}N$	$175^{\circ}E$	1200	500-4000	April-October	Kaneyasu and Murayami [2000]
Annual Antarctical	Ny-Alesund, Arctic	78°54'N	11°53′E	1900	516		Beine et al. [2001]
Mand Land, Antaretiea         73°3′S         13°2′S'W         30         228         December – February outs Island           Josts Island         43°S         119′T         249         1960         Ammal outs Ammal           Josts Island         43°S         119′T         249         1960         Ammal outs Ammal           Jost Island         43°S         119′T         260–3570         April Ammal outs Ammal           Islandic         13°N         60°2N         260         370         April Ammal outs Ammal           Islandic         47°N         84°W         88'O         2950         April Am           Islandic         27°N         81°W         830         2950         April Am           Islandic         27°N         81°W         830         2950         April Am           Islandic         27°N         81°W         380         2950         April Am           Islandic         27°N         48°N         3750         Ammal           Islandic         38°N         170         48°N         Ammal           Islandic         47°N         14°E         93         3760         Ammal           Islandic         48°N         14°E         176         Ammal </td <td>Palmer. Antarctica</td> <td>S°59</td> <td>64 °W</td> <td>45</td> <td>67</td> <td>Annual</td> <td>Savoie et al. [1993]</td>	Palmer. Antarctica	S°59	64 °W	45	67	Annual	Savoie et al. [1993]
volus Island         33°9 N         119°18 W         2490         1960         Annual Annual Annual           in Arch Atlantic         13°3 N         119°18 W         249         1960         Annual April           In North Atlantic         13°16 N         73°W         2670         260°370         Annual April           In North Atlantic         31°N         84°W         38°N         2570         Annual April           In Ontario         47°N         84°W         38°N         318         April           In Ontario         47°N         84°W         38°N         Annual April           In Ontario         47°N         81°W         38°N         Annual April           In Sweden         28°*48°N         17°23°W         48°N         Annual April           In Sweden         28°*48°N         17°23°W         48°N         Annual April           In Sweden         28°*48°N         17°23°W         48°N         Annual April           In Sweden         28°*48°N         14°N         48°N         48°N         Annual April           In Sweden         28°*8         14°N         14°N         48°N         Annual April           In Sweden         48°N         14°N         20°N	Oneen Mand Land Antarctica	2,2,2,2	13°25′W	30	228	December February	Kerminen et al [2000]
total statutu         1.3° s. 1.44° s. 1.9° s.	Con Misoles Island	12,00,00	110°1°W	2400	1060	Ammol	$V_{im} = A - C - C$
North Atlantic   13-167-N   60-137-N   143-E   250   260-3670   Application   13-167-N   143-E   250   260-3670   Application   13-167-N   143-E   250   260-3670   Application   13-18N   13-	Sali Inicolas Islaila	13 % IS	119 18 W	0647	155	Allinai	Nim et al. [2000]
17   17   17   17   17   17   17   17	Tasmania Tassis et Mende Adlende	45 S	144 E	230	133	December	berresneim ei di. [1990]
tine         37 M         75 M         2670         3200         January - December           tine         47 N         84 W         3870         470         Annual           tine         3 N         69 W         850         315         April - May           tine         27 N         81 W         850         Annual         Annual           tine         27 N         1723 W         1460         1400         June - July           ose, N. Y.         44 N         175 W         2950         Annual           ose, N. Y.         44 N         175 W         2120         2190         Annual           ose, N. Y.         44 N         147 M         147 M         Annual           ose, N. Y.         44 N         147 M         148 M         Annual           ose, N. Y.         44 N         147 M         148 M         Annual           ose, N. Y.         44 N         147 M         148 M         Annual           ose, N. Y.         48 N         147 M         148 M         Annual           ose, N. Y.         48 N         107 M         148 M         Annual           ose, N. Y.         48 N         107 M         140 M         Annual <td>Iropical North Atlantic</td> <td>13-16/-N</td> <td>60-23-W</td> <td>1620</td> <td>760-36/0</td> <td>April</td> <td>Johansen et al. [2000]</td>	Iropical North Atlantic	13-16/-N	60-23-W	1620	760-36/0	April	Johansen et al. [2000]
t, Ontario         47°N         84°W         3870         Annual           1         3°N         69°W         3870         Annual           1         27°N         81°W         3380         2950         Annual           1         27°N         17°24         1400         July - August           1         27°N         17°24         1460         July - August           1         44°N         75°W         1460         July - August           1         44°N         75°W         1400         June - July           1         50°N         75°W         1400         June - July           1         44°N         75°W         2120         2190         Annual           1         50°N         75°W         2120         2190         Annual           1         44°N         75°W         2120         2190         Annual           1         44°N         14°W         2120         2190         Annual           1         44°N         16°W         220         2100         Annual           2         48°N         10°W         2040         940         Annual           3         46°N         <	West Atlantic	31°N	75°W	2670	3200	January – December	Berresheim et al. [1990]
t, Ontario         47°N         84°W         3870         2570         Amual           1, Ontario         3°N         69°W         3870         315         April—May           1, Ontario         3°N         870         2570         April—May           1d, Fla.         27°N         81°W         3380         2950         Amual           en, Sweden         58°4K         17°34W         1460         1400         Ime—luly           en, Sweden         58°4K         17°34W         2120         2190         Amual           en, N. Y.         44°N         75°W         2120         2190         Amual           sys. S. D.         50°N         14°E         3340         1765         Amual           s, Quebec         4°N         14°E         935         835         June           s, Quebec         4°N         107°W         2040         940         Amual           s, Quebec         4°N         107°W         2040         940         Amual           sk, Quebec         4°N         107°W         2040         940         Amual           sk, Quebec         4°N         19°4         45°N         140°N         140°N         140°N <td></td> <td></td> <td></td> <td>Rural</td> <td></td> <td></td> <td></td>				Rural			
def. Fla.         3°N         69°W         850         315         April-May           dd, Fla.         27°N         81°W         350         2485         July-August           en, Sweden         58°48°N         17°23′W         1460         1400         July-August           en, Sweden         58°48°N         17°23′W         1460         1400         July-August           ose, N. Y.         44°N         75°W         4830         2350         July-August           ose, N. Y.         44°N         75°W         4830         2120         2190         Ammal           ose, N. Y.         44°N         75°W         3340         1765         Ammal           is, Oubecc         50°N         14°E         935         835         June           ke, Saskarchewan         57°N         107°W         2040         940         Ammal           ke, Minn.         48°N         107°W         2600         940         Ammal           ke, Minn.         48°N         19°42         4550         2670         July-August           ke, Minn.         46°57         19°42         4550         2700         400         Ammal           st Greenland         47°5	Algoma, Ontario	N°74	84°W		2570	Annual	McNauohton and Vet [1996]
Id. Fla.         27°N         81°W         1200         48.5         July-Trianger           ee, Sweden         58°48'N         17°23'W         1460         1400         Jule-July           ee, Sweden         58°48'N         17°23'W         1460         1400         June-July           ee, Sweden         58°48'N         17°24'N         1460         Jule-July         Amual           igs, S. D.         44°N         57°W         2120         2190         Amual           is, Quebec         50°N         17°W         3340         1765         Amual           ke, Saskatchewan         57°N         107°W         2040         Amual           ke, Saskatchewan         57°N         107°W         2040         Amual           g. Minn.         4°N         58°W         2040         Amual           g. Minn.         4°N         58°W         200         1385         Amual           ski, Russia         69°N         29°E         5870         July-August           ski, Russia         69°N         29°E         4550         260         1710         Amual           ski, Russia         69°N         46°S         46°S         46°S         46°S         46°	Amazon	No.	/M <sub>0</sub> 09	058	315	April_Max	Talbat at al [1990]
Id, Fla.         27°N         81°W         1200         76°J         Januar Anguar           en, Sweden         58°48°N         17°23°W         1460         1400         Januar Anguar           ose, N. Y.         44°N         17°23°W         1460         1400         Januar July           dose, N. Y.         44°N         17°24         1460         1400         Januar July           dos, N. Y.         44°N         57°W         2120         2190         Amual           de, S. D.         44°N         14°E         935         835         June           de, Saskatchewan         57°N         107°W         2640         940         Amual           g, Minn.         48°N         91°W         2600         940         Amual           g, Minn.         48°N         91°W         2600         170         Amual           g, Minn.         48°N         10°W         260         250         July         Amual           g, Min.         49°N         18°40         45°0         1400         1400         Amual           g, Was         44°N         16°40°W         1100         490         Amual           s, South Africa         24°30°N	MIIIAZOII	AT C	*	1200	515	Tapin-indy Indy Amengt	10001 et al. [1990]
Affice and September 1, September 1, September 2, September	-	i d	1330 0 0	1200	000	Juny-August	[] () () () () () () () () () () () () ()
ren, Sweden         58°48°N         17°23°W         1460         1400         June-July           ose, N. Y.         49°N         17°23°W         4830         3750         Amual           ose, N. Y.         44°N         57°W         4830         3750         Amual           is, Quebec         50°N         75°W         3340         1765         Amual           is, Quebec         50°N         14°E         935         835         June           is, Quebec         49°N         107°W         2040         1760         Amual           is, Quebec         50°N         107°W         2040         October         October           is, Quebec         50°N         107°W         2040         October         October           is, Quebec         50°N         20°N         200         October         October           is, Quebec         50°N         20°N         200         October         October           is, Quebec         58°W         200         250         Amual           is, Quebec         58°W         20°         250         July-August           is, Rissia         69°N         140°N         140°         Amual	Archbold, Fla.	Z/2Z	8] W	3380	2950	Annual	McNaughton and Vet [1996]
Ose, N. Y.         44°N         75°W         4830         3750         Anmual Anmual Anmual Anmual Anmual Anmual Anmual Andual Anmual Anmual Anmual Anmual Anmual Anmual Anmual Anmual Andual Anmual Anm	Aspvreten, Sweden	58°48'N	17°23′W	1460	1400	June-July	Zappoli et al. [1999]
gs, S. D.         44°N         97°W         2120         Annual           5, Quebec         50°N         75°W         3340         1765         Annual           4°N         14°E         935         835         June           ke, Saskatchewan         57°N         107°W         2040         940         Annual           g, Minn.         48°N         91°W         2040         Annual           g, Minn.         48°N         91°W         2040         Annual           g, Minn.         48°N         50°         250         Annual           g, Minn.         48°N         29°E         5870         Annual           ski, Russia         69°N         28°E         4550         L40°-12,600         July-August           a, Hungary         46°57'N         19°42'E         4550         2670         July-August           dujik, Nova Scotia         47°N         56°W         3550         2870         July-August           st Greenland         81°36'N         16°40'W         1100         490         Annual           st Greenland         24°38'N         16°40'W         1100         490         Annual           o., Wis.         45°N         89°W </td <td>Big Moose, N. Y.</td> <td>4 N°</td> <td>75°W</td> <td>4830</td> <td>3750</td> <td>Annual</td> <td>McNaughton and Vet [1996]</td>	Big Moose, N. Y.	4 N°	75°W	4830	3750	Annual	McNaughton and Vet [1996]
s, Quebec         50°N         75°W         3340         1765         Annual           4°N         14°E         935         835         June           1040         0ctober         1040         October           1170         107°W         2040         940         Annual           g, Minn.         4°N         58°W         2600         1385         Annual           ski, Russia         69°N         29°E         5870         June         Annual           ski, Russia         69°N         29°E         5870         July-August           4°St, Russia         46°STN         19°42'E         4550         July-August           46°STN         19°42'E         4550         2670         July-August           46°STN         19°42'E         4550         2670         July-August           46°SY         50°W         3550         2840         Annual           sst Greenland         4°S'N         16°40'W         1100         A90         Annual           st Greenland         4°S'N         16°40'W         1100         490         Annual           st South Africa         28°24'E         2650         5890-13,800         Annual	Brookings, S. D.	N°4	M <sub>0</sub> L6	2120	2190	Annual	McNaughton and Vet [1996]
ke, Saskatchewan         4°N         14°E         935         835         June           ke, Saskatchewan         57°N         107°W         2040         940         Annual           g, Minn.         48°N         91°W         260         1385         Annual           g, Minn.         48°N         58°W         920         250         June           ski, Russia         69°N         29°E         5870         July         Annual           a, Hungary         46°57'N         19°42'E         4550         2670         July-August           a, Koa         46°57'N         16°40'W         3350         2840         Annual           ss         4°58'N         16°40'W         1100         490         Annual           sst Greenland         4°58'N         16°40'W         3690         5800         490         Annual           o, Wis.         45°N <t< td=""><td>Chapais, Quebec</td><td><math>N_{\circ}0S</math></td><td><math>^{2}</math></td><td>3340</td><td>1765</td><td>Annual</td><td>McNaughton and Vet [1996]</td></t<>	Chapais, Quebec	$N_{\circ}0S$	$^{2}$	3340	1765	Annual	McNaughton and Vet [1996]
ke, Saskatchewan         57°N         107°W         107°W         2040         Amual           g, Minn.         4°N         58°W         2040         940         Amual           sk, Minn.         4°N         58°W         2040         1385         Amual           ski, Russia         69°N         29°E         570         1710         Amual           a, Hungary         46°57'N         19°42'E         4550         2670         July-August           46°57'N         19°42'E         4550         1400-12,600         July-August           46°57'N         19°42'E         4560         5760         July-August           46°57'N         19°42'E         4550         1400-12,600         July-August           47°0         77°0         2700-9400         February           st Greenland         44°N         16°40'W         1100         490         Amual           st Greenland         24°30'N         28°4'E         2650         5890-13,800         Amual           oo, Wis:         45°N         89°W         3690         3120         Amual           Colo.         40°N         107°W         1760         1080         Amual	Congo	$^{6}$	14°E	935	835	June	Clairac et al. [1988]
57°N         107°W         2040         940         Annual           48°N         91°W         2600         1385         Annual           4°N         58°W         920         250         June           69°N         29°E         5870         1710         Annual           46°57N         19°42°E         4550         2670         July-August           46°57N         19°42°E         4550         July-August           46°57N         4500         5760         Annual           47°N         3550         2840         Annual           47°N         1100         280         Annual           24°36'N         1100         490         Annual           45°N         80°W         3690         3120         Annual           45°N         69°W         3690         3120         Annual           40°N         107°W         1760         1080         Annual				1170	1040	October	
48°N         91°W         2600         1385         Annual           4°N         58°W         920         250         June           69°N         29°E         5870         1710         Annual           46°57'N         19°42'E         4550         2670         July-August           40°N         45°0         1400-12,600         July-August           40°N         55°W         350         2840         Annual           4°58'N         73°28'E         770         2700-9400         February           81°36'N         16°40'W         1100         490         Annual           45°N         89°W         3090         3410         Annual           45°N         69°W         3690         3410         Annual           40°N         10°W         1760         1080         Annual	Cree Lake, Saskatchewan	$N^{\circ}78$	$107^{\circ}$ M	2040	940	Annual	McNaughton and Vet [1996]
4°N         58°W         920         250         June           69°N         29°E         5870         1710         Annual           46°S7'N         19°42'E         4550         2670         July-August           400         2760         July-August           400         5760         Annual           Annual         5760         Annual           Annual         770         2700-9400         February           81°36'N         16°40'W         1100         490         Annual           45°N         89°W         3090         3410         Annual           45°N         69°W         3690         3120         Annual           40°N         107°W         1760         1080         Annual	Fernberg, Minn.	$^{48}^{\circ}N$	$M_{\circ}16$	2600	1385	Annual	McNaughton and Vet [1996]
69°N         29°E         5870         1710         Annual           46°57'N         19°42'E         4550         2670         July-August           46°57'N         19°42'E         4550         1400-12,600         July-August           4000         5760         Annual           Annual         5760         Annual           A°58'N         73°28'E         770         2700-9400         February           81°36'N         16°40'W         1100         490         Annual           45°N         89°W         3090         3410         Annual           45°N         69°W         3690         3120         Annual           40°N         107°W         1760         1080         Annual	Guyana	$^{\circ}$ V	28°W	920	250	June	Gregory et al. [1986]
46°57'N         19°42'E         4550         2670         July-August           46°57'N         19°42'E         4550         1400-12,600         July-August           4000         5760         Annual           4000         5760         Annual           Annual         Annual         Annual           4°58'N         73°28'E         770         2700-9400         February           81°36'N         16°40'W         1100         490         Annual           45°N         89°W         3090         3410         Annual           45°N         69°W         3690         3120         Annual           40°N         107°W         1760         1080         Annual	Janiskoski, Russia	N <sub>0</sub> 69	29°E	5870	1710	Annual	Hjellbrekke and Hanssen [1998]
4550         1400–12,600         July-August           4000         5760         Annual           4000         5760         Annual           4000         5760         Annual           4°58'N         770         2840         Annual           81°36'N         16°40'W         1100         490         Annual           24°39'N         28°24'E         2650         5890–13,80         Annual           45°N         89°W         3090         3120         Annual           45°N         69°W         3690         3120         Annual           40°N         107°W         1760         1080         Annual	K-puszta, Hungary	46°57′N	19°42′E	4550	2670	July-August	<i>Temesi et al.</i> [2001]
ia         44°N         65°W         3550         5760         Annual           4°58'N         73°28'E         770         2700–9400         February           81°36'N         16°40'W         1100         490         Annual           24°39'N         28°24'E         2650         5890–13,800         Annual           45°N         89°W         3090         3410         Annual           45°N         69°W         3690         3120         Annual           40°N         10°W         1760         1080         Annual				4550	1400 - 12.600	July-August	Zappoli et al. [1999]
ia 44°N 65°W 3550 2840 Annual 4°58'N 73°28'E 770 2700–9400 February 81°36'N 16°40'W 1100 490 Annual 24°39'N 28°24'E 2650 5890–13,800 May 45°N 89°W 3090 3120 Annual 45°N 69°W 3690 3120 Annual 40°N 107°W 1760 1080 Annual				4000	5760	Annual	Hiellbrekke and Hanssen [1998]
4°58N 73°28E 770 2700–9400 February 1100 490 Annual 24°39N 28°24'E 2650 5890–13,800 May 45°N 89°W 3090 3120 Annual 45°N 69°W 3690 1760 1080 Annual Annual 40°N 107°W 1760 1080 Annual	Keiimkniik Nova Scotia	N°44	M°59	3550	2840	Annual	McNauohton and Vet [1996]
81°36′N 16°40′W 1100 490 Annual 24°39′N 28°24′E 2650 5890–13,800 May 45°N 89°W 3090 3120 Annual 40°N 10°0′W 1760 1080 Annual	Maldives	N/85°L	73°78'F	022	0700-0400	February	Chowdhum of al [2001]
24.39'N 28°24'E 2560 5890–13,800 Maay 45°N 89°W 3090 3120 Annual 40°N 107°W 1760 1080 Annual	Northeon Greenland	01026'N	16°40'W	1100	400	Annual	Document = 10001
24.39 N 28.24 E 2650 3890-15,800 May 45°N 89°W 3690 3410 Annual 45°N 69°W 3690 3120 Annual 40°N 1760 1080 Annual	Notificast Officiality	01 30 14	10 +0 W	0011	064	Aminai	11etaam et at. [1998]
45°N 89°W 3090 3410 Annual Annual Annual 45°N 69°W 3690 3120 Annual 40°N 107°W 1760 1080 Annual	Nyisviey, South Africa	24-39'N	28-24 E	2650	5890-13,800	May	Fuxbaum et al. [2000]
45°N 69°W 3690 3120 Annual 40°N 107°W 1760 1080 Annual	Shawano, Wis.	N <sup>5</sup> 24	W-68	3090	3410	Annual	McNaughton and Vet [1996]
40°N 107°W 1760 1080 Annual	Winterport, Maine	45°N	M <sub>2</sub> 69	3690	3120	Annual	McNaughton and Vet [1996]
	Yampa, Colo.	V°04	$107^{\circ}$ W	1760	1080	Annual	McNaughton and Vet [1996]

Station	Latitude	Longitude	Modeled sulfate $(ng/m^3)$	Measured sulfate (ng/m³)	Period	Data reference
			Urban			
Arendtsville, Pa.	$40^{\circ}N$	$M_{\circ}LL$	4600	6773	Annual	McNaughton and Vet [1996]
Atlanta, Ga.	$33^{\circ}N$	M°28	3500	2000-20,500	August	Modey et al. [2001]
Beijing, China	$^{40}$ N	116°E	3630	14,250	Annual	He et al. [2001]
			2790	17,140	Summer	
			3240	12,550	Autumn	
			4330	24,870	Winter	
			4000	10,150	Spring	
Bells, Tenn.	$36^{\circ}N$	$M_{\circ}68$	4740	5170	Annual	McNaughton and Vet [1996]
Bilthoven, Netherlands	$52^{\circ}N$	$5^{\circ}\mathrm{E}$	3630	4400	Annual	Hjellbrekke and Hanssen [1998]
Due West, S. C.	$34^{\circ}N$	$M_{\circ}$ 8	3900	5840	Annual	McNaughton and Vet [1996]
Fort Wayne, Indiana	$41^{\circ}N$	%58 Wo	5730	6300	Annual	McNaughton and Vet [1996]
Ispra, Italy	$46^{\circ}N$	$_{0}^{\circ}$ E	2860	5600	Annual	Hjellbrekke and Hanssen [1998]
Ithaca, N. Y.	$42^{\circ}N$	$M_{\circ}9L$	5070	5000	Annual	McNaughton and Vet [1996]
Jarczew, Poland	$51^{\circ}N$	$22^{\circ}E$	4910	9130	Annual	Hjellbrekke and Hanssen [1998]
Jerome, Miss.	$38^{\circ}N$	$M_{\circ}$ Z6	3910	4130	Annual	McNaughton and Vet [1996]
Kosetice, Czech Republic	$20^{\circ}N$	$15^{\circ}$ E	3990	0969	Annual	Hjellbrekke and Hanssen [1998]
La Cartuja, Spain	$37^{\circ}N$	$^{\circ}$ M	2170	4020	Annual	Hjellbrekke and Hanssen [1998]
Logrono, Spain	$42^{\circ}N$	$^{\circ}M$	2290	3290	Annual	Hjellbrekke and Hanssen [1998]
Los Angeles Basin	34°4′N	118°15′W	2330	4240 - 5190	Annual	Kim et al. [2000]
Marion, Ala.	$33^{\circ}N$	$M_{\circ}$	4380	5190	Annual	McNaughton and Vet [1996]
Marshall, Tex.	$33^{\circ}N$	94°W	2760	5110	Annual	McNaughton and Vet [1996]
Mountain Lake, Va.	$37^{\circ}N$	$81^{\circ}$ W	3790	6710	Annual	McNaughton and Vet [1996]
Pittsboro, N. C.	36°N	M <sub>0</sub> 6L	3800	6041	Annual	McNaughton and Vet [1996]
Po Valley, Italy	44°39′N	$11^{\circ}37/E$	3150	3400 - 11,000	September	Zappoli et al. [1999]
Roquetas, Spain	$41^{\circ}N$	$1^{\circ}W$	2340	3750	Annual	Hjellbrekke and Hanssen [1998]
Suwalki, Poland	$54^{\circ}N$	23°E	5730	6220	Annual	Hjellbrekke and Hanssen [1998]
Svratouch, Czech Republic	$20^{\circ}N$	$16^{\circ}\mathrm{E}$	4160	5710	Annual	Hjellbrekke and Hanssen [1998]
Thessaloniki	40°31′N	22°58′E	4250	4500 - 8200	June	Chazette and Liousse [2001]
Toledo, Spain	$^{40}$ N	$4^{\circ}$ W	2200	1820	Annual	Hjellbrekke and Hanssen [1998]
Uvalda, Ga.	$32^{\circ}N$	82°W	3910	4470	Annual	McNaughton and Vet [1996]
Vysokoe, Belarus	$52^{\circ}N$	23°E	5150	3960	Annual	Hjellbrekke and Hanssen [1998]
Warwick, Mass.	$43^{\circ}N$	$72^{\circ}$ W	4680	4360	Annual	McNaughton and Vet [1996]
Witteveen, Netherlands	$53^{\circ}N$	2°E	3980	3780	Annual	Hjellbrekke and Hanssen [1998]
Zanesville, Ohio	$40^{\circ}N$	85°W	4880	0869	Annual	McNaughton and Vet [1996]
						Ì







**Figure 4.** Comparison of yearly and zonally averaged modeled with (a) satellite-derived cloud water [*Greenwald et al.*, 1995], (b) climatological precipitation [*Xie and Arkin*, 1996], and (c) Nimbus-7 satellite cloud fraction data [*Hack et al.*, 1998].

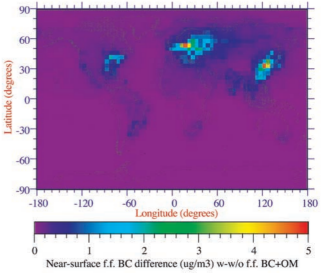
[70] Comparisons of model predictions of direct forcing with those from other studies are given by Jacobson [2001b] for several components. Of relevance, the direct forcing per unit mass of organic carbon (OC) (= OM/1.3) was  $-64 \text{ W/m}^2/\text{g}$  OC when UV absorption by organics was ignored, consistent with  $-70 \text{ W/m}^2/\text{g}$  OC from the work of  $Cooke\ et\ al.$  [1999]. When UV absorption by organics was accounted for, the OC direct forcing was  $-43 \text{ W/m}^2/\text{g}$  OC.

[71] Another source of uncertainty is cloud and precipitation fields. Figures 4a–4c compare yearly and zonally averaged cloud water, precipitation, and cloud fraction with satellite-derived or climatological fields. In all cases, the model predicted observed general characteristics. Due in part to the coarse model resolution, much uncertainty still exists in predicting the location, height, water content, and optical characteristics of clouds.

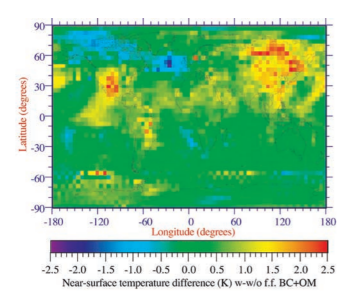
# 6. Analysis of Climate Response and Individual Effects

[72] Figures 5–14 are difference plots for several variables, all obtained by taking the difference in results in year 5 when f.f. BC + OM emissions were and were not included, respectively. In both cases, BC and all other emissions from biomass burning were included. The only difference between the two simulations was f.f. BC + OM emissions.

[73] Figure 5 shows the difference in near-surface f.f. BC from the two simulations. Most f.f. BC originated from the eastern United States, Europe, eastern Asia, and parts of South America and Africa. Figure 6 shows the difference in year 5 in near-surface temperatures resulting from the f.f. BC + OM simulation. The globally averaged difference was +0.35 K. Most the globe warmed due to BC + OM, but not necessarily in the regions where near-surface BC + OM concentrations were highest. The major regions of warming were over Eastern Russia, Southeast Asia, and Southwestern North America. The major region of cooling was over the North Atlantic. Both the regions of warming and cooling are consistent in location with major observed regions of warming and cooling, respectively, between 1949 and 1997 [Knutson et al., 1999, Plate 3a; Schneider and Held, 2001, Figure 1a + c]. The results here account for some of the observed trends even when perturbations to greenhouse gases are not considered, implying that a portion of such observed trends may be due to f.f. BC + OM.



**Figure 5.** (a) Fifth-year average difference in near-surface f.f. BC concentration ( $\mu g/m^3$ ) when f.f. BC + OM emissions were versus were not included in the calculation. (b) Same as (a), but for zonally averaged BC ( $\mu g/m^3$ ).



**Figure 6.** Same as Figure 5a, but for near-surface air temperature (K).

[74] Figure 7a shows the globally and time-averaged layer and cumulative temperature differences after the transient f.f. BC + OM and equilibrium f.f. CO<sub>2</sub> removal simulations. Whereas, f.f. CO<sub>2</sub> and f.f. BC + OM warmed the lower troposphere, CO<sub>2</sub> cooled the upper troposphere and stratosphere (because absorption of thermal-IR by CO<sub>2</sub> in the lower troposphere prevented that thermal-IR from being absorbed in the stratosphere). The maximum cooling from 30 to 100 mb was 1.5 K, which compares with a 1.5-2 K cooling from radiosonde measurements [Angell, 1999] due to all climate perturbations at 30-100 mb since 1958. As such, a significant portion of stratospheric cooling since 1958 appears to be due to greenhouse gas buildup in the troposphere. The large modeled decrease in upper stratospheric temperatures found here is supported by Akmaev and Fomichev [2000], who used a global model to predict a cooling of 3 K at 50 km when CO<sub>2</sub> was increased from 313 to 360 ppmv. Scaling their cooling by the ratio of the CO<sub>2</sub> change assumed here (280 to 365 ppmv) to that assumed there gives a cooling from their results of 5.4 K, which compares with a cooling here of 5.8 K.

[75] The f.f. BC + OM increased air temperatures relative to ground temperatures (Figure 7a), increasing the stability of near-surface air, slightly slowing near-surface winds, reducing low troposphere and midtroposphere cumulus convection (Figure 7b), increasing low-cloud liquid water (Figure 7c), decreasing middle cloud liquid water and ice and upper cloud ice (Figure 7c), and reducing precipitation (Table 3). Despite overall lower cloud liquid water and ice contents (Table 3 and Figure 7c), the "first indirect effect" increased the 550 nm cloud scattering optical depth at these levels (Figure 7d). Because BC existed within, above, and below cloud layers (e.g., Figure 7e), the enhanced cloud scattering increased solar plus thermal-IR heating within and above clouds up to 250 mb (Figure 7b).

[76] The warming due to BC within the boundary layer and within and above cloud layers increased the downward sensible heat flux to the surface, increasing surface temperatures although not to the same extent as atmospheric

temperatures. For example, f.f. BC + OM increased surface temperatures by about 0.28 K, which was 0.07 K less than near-surface air temperatures increased in the global average (Figure 7a). Due to the increases in surface temperatures, more ocean and soil water evaporated, increasing water vapor contents (Figure 7d). Increases in tropospheric and lower stratospheric water vapor is an observed trend [Oltmans and Hofmann, 1995]. Since the air was warmer, saturation vapor pressures were higher, allowing more water vapor to accumulate and reducing cloud liquid, cloud ice, and precipitation in the global average (Table 3). The accumulation of water vapor increased solar-IR and thermal-IR heating rates, increasing air temperatures further ("BC-water vapor positive feedback").

[77] Figure 7e shows the change in globally averaged vertical profiles of BC, S(VI), Na<sup>+</sup>, and soil dust resulting when f.f. BC + OM was emitted. The figure shows that f.f. BC + OM increased sodium and sulfate but decreased soil dust. Sulfate and sodium increased partly because precipitation decreased over the oceans and over northern latitudes, where these species were abundant ("rainout effect"). Soil dust decreased, possibly due to the "daytime stability" and "smudge-pot effects" and to the fact that precipitation did not decrease so much over soil dust source regions (deserts). The figure also shows that increases in BC and S(VI) concentrations exceeded those at the surface, suggesting that these components were removed more readily near the surface than aloft by rainout, dry deposition, and sedimentation. For example, rainout from low-lying stratus producing drizzle diminish aerosols near the surface but not above the stratus.

[78] Figure 7g shows the difference in downward minus upward solar, thermal-IR, and total irradiance (total forcing) from the f.f. BC + OM simulations. The solar irradiance difference was positive at the tropopause and negative at the surface, indicating in part that BC and the additional water vapor it produced absorbed solar and solar near-IR radiation, respectively. The reduced downward (increased upward) thermal-IR forcing at the tropopause resulted in part because f.f. BC + OM reduced middle and upper tropospheric cloud liquid water and ice, reducing the downward flux of thermal-IR because liquid water and ice are thermal-IR absorbers. Downward thermal-IR irradiance decreased less at lower altitudes, in part because water vapor and cloud liquid contents increased most down there.

[79] The positive net solar plus thermal-IR tropopause forcing due to f.f. BC + OM is partly explained as follows. BC absorbed sunlight, heating the air (positive forcing). BC + OM increased water vapor, enhancing positive forcing more. BC + OM increased the number of CCN, increasing negative forcing through the "indirect effects", but it reduced middle and upper (and overall) cloud liquid water and ice contents, offsetting some of this negative forcing. The enhanced scattering due to the "first indirect effect" enhanced absorption by BC, H<sub>2</sub>O, and CO<sub>2</sub>, within and above clouds, increasing positive forcing. BC + OM also reduced surface albedo (Table 3), increasing positive forcing.

[80] A comparison of the positive net forcing due to f.f. BC + OM with estimates of negative "indirect" forcing is a comparison of apples and oranges. The net forcing here is due to direct forcing plus a multitude of time-dependent

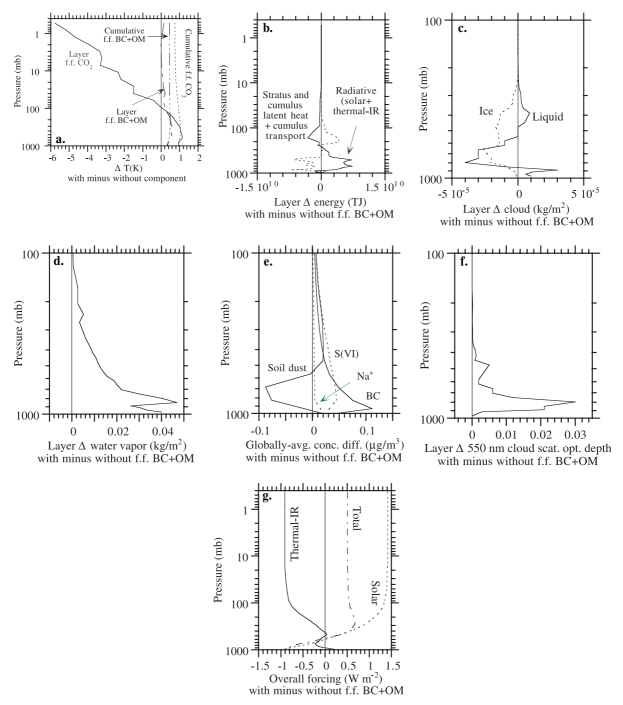


Figure 7. (a) Difference in globally and time-averaged layer and cumulative temperatures when all aerosols and gases were present versus when f.f. BC + OM emissions were removed (transient simulation) and anthropogenic CO<sub>2</sub> above 280 ppmv was removed (equilibrium simulation). The globally and time-averaged temperature of a layer is  $T_{layer} = \sum_{c,t} V_{c,t} \rho_{a,c,t} c_{p,a} T_{c,t} / \sum_{c} V_{c,t} \rho_{a,c,t} c_{p,a}$ , where the summation is over all grid cells in the layer and over all time steps during the averaging period,  $V_{c,t}$  is the volume of a grid cell at a given time step,  $\rho_{a,c,t}$  is the air density in a grid cell at a given time step,  $c_{p,a}$ is the specific heat of air at constant pressure, and  $T_{c,t}$  is the temperature in a grid cell at a given time step. The cumulative temperature uses a similar equation, except that the integration includes all grid cells from the ground to the altitude of interest. Thus, the cumulative temperature at the stratopause is the integration of air temperature from the surface to the stratopause. The bottom-most "layer" temperature changes are ground-temperature changes, which are not included in the cumulative temperature change curves. The remaining curves are global differences from the simulations with minus without f.f. BC + OM for (a) changes in energy due to latent heat release plus cumulus energy transport and solar plus thermal-IR heating, (b) cloud liquid and ice, (c) cloud scattering optical depth, (d) several aerosol constituents, (e) water vapor, and (f) net downward minus upward solar, thermal-IR, and total irradiance.

Parameter	Difference	Parameter	Difference
Near-surface temperature	+0.35 K	Near-surface aerosol single-scattering albedo	-0.029 (-)
Near-surface relative humidity	-0.0017 (out of 1)	Near-surface aerosol LWC	$+0.17 \mu g/m^3$
Surface pressure	-0.0003 mb	Near-surface BC	$+0.11  \mu g/m^3$
Surface albedo	-0.0017 (out of 1)	Near-surface OM	$+0.23  \mu \text{g/m}^3$
Cloud optical depth	+0.27 (-)	Near-surface S(VI)	$+0.052 \mu \text{g/m}^3$
Cloud liquid water	$-0.000089 \text{ kg/m}^2$	Near-surface NO <sub>3</sub>	$+0.003  \mu \text{g/m}^3$
Cloud ice	$-0.00019 \text{ kg/m}^2$	Near-surface Cl	$+0.004  \mu \text{g/m}^3$
Precipitation	-0.017  mm/day	Near-surface NH <sub>4</sub> <sup>+</sup>	$+0.011  \mu \text{g/m}^3$
Tropopause total sol + IR forcing	$+0.53 \text{ W/m}^2$	Near-surface Na <sup>+</sup>	$+0.0025 \mu g/m^3$
Aerosol optical depth	+0.023 (-)	Near-surface SO <sub>2</sub>	+0.0089 ppbv

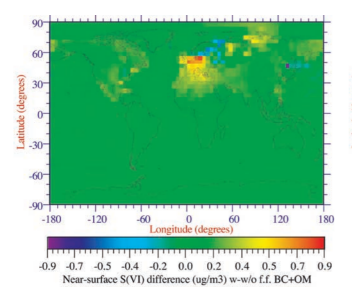
**Table 3.** Fifth-Year Globally Averaged Difference in Several Parameters Resulting From the Simulations When f.f. BC + OM Emissions Were Versus Were Not Included

effects, including indirect effects, when all effects are linked together. Estimates of indirect forcing, which range from 0 to  $-2 \text{ W/m}^2$  [IPCC, 2001] are based on simulations in which forcing is ascribed to indirect effects only and not to direct effects of f.f. BC + OM nor the effects of the multitude of feedbacks treated here. Nevertheless, the facts that (1) cloud scattering optical depths increased at all altitudes when f.f. BC + OM was included (Figure 7d) and (2) that the inclusion of f.f. BC + OM reduced the mean cloud radius in the global average suggest that indirect forcing here due to f.f. BC + OM was negative. In addition, the total forcing obtained for the with/without f.f. SO<sub>2</sub> simulations was negative, indicating again that indirect forcing must have been negative. It is impossible to isolate the magnitude of indirect forcing without holding many parameters constant, thereby changing the climate responses and rendering the results incorrect (section 3.13), so this was not attempted here.

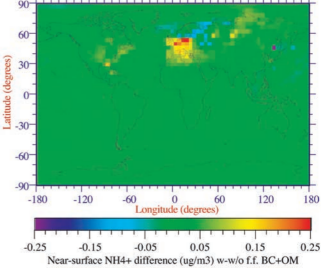
[81] Figure 8 shows the globally distributed change in near-surface S(VI)(aq) due to the emission of f.f. BC + OM. The increase in particle surface area over western Europe due to f.f. BC + OM appears to have enhanced condensation of H<sub>2</sub>SO<sub>4</sub>(g), increasing aerosol liquid water content,

ammonia uptake (Figure 9), and aqueous oxidation of SO<sub>2</sub> to S(VI)(aq) (all "self-feedback effects"). Because more S(VI)(aq) was produced in western Europe, less was produced in northeastern Europe (Figure 8). At the same time, precipitation decreased over western Europe, decreasing the removal of S(VI)(aq) and SO<sub>2</sub> ("rainout effect"), so it is not clear the extent to which the enhanced S(VI)(aq) over western Europe was due to the "self-feedback effect" versus the "rainout effect."

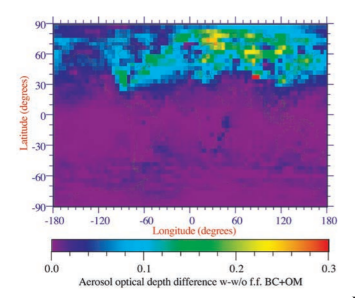
[82] Figure 10 shows the change in aerosol optical depth (integrated from the surface to top-of-atmosphere) due to the emission of f.f. BC + OM. Aerosol optical depth increased particularly in the Northern Hemisphere, not only due to the addition of f.f. BC + OM but also due to resulting increases in S(VI)(aq), NO<sub>3</sub><sup>-</sup>, Cl<sup>-</sup>, NH<sub>4</sub><sup>+</sup>, Na<sup>+</sup>, and aerosol liquid water (Table 3). The greater number concentration of particles (mainly f.f. BC + OM and homogeneously nucleated S(VI)(aq), decreased effective cloud radii and increased cloud optical depths, particularly over Europe and parts of Asia (Figure 11). The change in cloud optical depths reflects the sum of "indirect effects", the "semi-direct effect", and the "BC-low-cloud positive feedback loop". The fact that cloud optical depths increased in the



**Figure 8.** Same as Figure 5a, but for near-surface aerosol S(VI) ( $\mu g/m^3$ ).



**Figure 9.** Same as Figure 5a, but for near-surface aerosol NH $_{+}^{4}$  (µg/m $^{3}$ ).



**Figure 10.** Same as Figure 5a, but for aerosol optical depth.

global average (Table 3) suggests that "indirect effects" may have dominated the other two.

[83] Figure 12 shows the change in surface pressures and winds. Changes in winds and pressures affected not only temperatures ("particle effect through large-scale meteorology"), but also clouds and precipitation (Table 3). Changes in winds also affected the emission rates of sea-spray and the Na<sup>+</sup> it contains (Figure 13). The change in Na<sup>+</sup> affected the liquid water content of such drops. For example, a decrease in sodium in the Norwegian Sea decreased aerosol liquid water content noticeably there.

[84] In the global average, the addition of f.f. BC + OM decreased surface albedo (Table 3) primarily due to a slight reduction in Northern Hemisphere land-snow cover (mostly over eastern Russia and Canada. The decreases over eastern Russia coincided with the increase in temperatures there, although it cannot be determined whether the change in

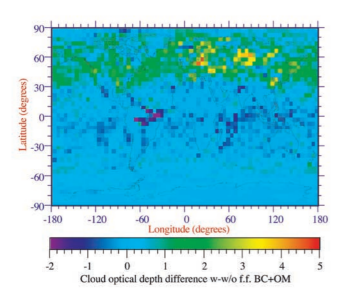
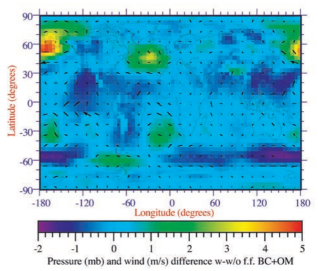


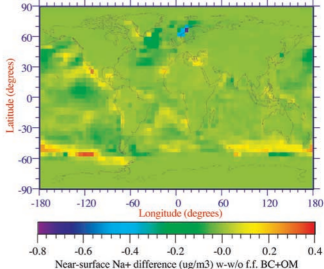
Figure 11. Same as Figure 5a, but for cloud optical depth.



**Figure 12.** Same as Figure 5a, but for surface pressure (mb) and winds (m/s).

albedo caused the change in temperature or vice versa or both.

[85] Finally, in the simulations in which f.f. SO<sub>2</sub> emissions were and were not included, the northeast United States, central and eastern Europe, and much of eastern Asia cooled. Warming occurred over much of the Southern Hemisphere, but was smaller than Northern Hemisphere cooling. A reason for the overall cooling was the effect of f.f. SO<sub>2</sub> emissions on cloud optical depth. Cloud optical depth increased when f.f. SO<sub>2</sub> emissions were included, because f.f. SO<sub>2</sub> produced sulfuric acid—water aerosol particles, which enhanced the "indirect effects". The change in cloud optical depth due to f.f. SO<sub>2</sub> was greater than that due to f.f. BC + OM, partly explaining why the magnitude of the climate response due to f.f. SO<sub>2</sub> was greater than that due to f.f. BC + OM. The cloud optical depth resulting from f.f.



**Figure 13.** Same as Figure 5a, but for near-surface aerosol Na<sup>+</sup> ( $\mu$ g/m<sup>3</sup>).

 $SO_2$  was greater than that due to f.f. BC + OM, because f.f.  $SO_2$ , spread further in the Northern Hemisphere than did f.f. BC + OM (e.g., Figure 3 versus Figure 2). As such, f.f.  $SO_2$  affected clouds over a greater global area than did f.f. BC + OM. The total (direct plus other) forcing resulting from the  $SO_2$  simulations was negative, indicating that the forcing due to the "indirect effects", if they could be isolated, must also have been negative.

# 7. Comparison of Diesel Versus Gasoline

[86] Two practical strategies to reduce BC + OM are to improve particle traps for diesel vehicles and to replace diesel with gasoline. A modern diesel direct injection (DDI) engine obtains 25–35% better mileage than an equivalent port fuel injection (PFI) gasoline engine, but much of this gain is offset, since diesel releases 18.1% more CO<sub>2</sub> per unit volume than does gasoline [Wang, 1999]. Although light-duty diesel engines release less CO<sub>2</sub> than do PFI gasoline engines, they release up to 0.08 g particulates/mi (0.05 g/km) under current U.S. and E.U. emission standards. Most particulate emissions are BC + OM [Shauer et al., 1999; Lighty et al., 2000]. New PFI engines produce 25 to >200 times less particulate mass per distance driven than new DDI engines [Maricq et al., 1999b].

[87] If a new PFI gasoline engine obtains 25 mpg (10.6) km/l), it releases about 95.4 g C/mi (59.4 g C/km) as CO<sub>2</sub> (with gasoline density of 737 g/l and carbon content of 85.5%) [Wang, 1999]. An equivalent DDI engine with 30% better mileage obtains 32.5 mpg (13.8 km/l), releasing 86.7 g C/mi (54.0 g C/km) as CO<sub>2</sub> (assuming diesel density of 856 g/l and carbon content of 87.0%) [Wang, 1999]. Thus, the DDI engine releases 8.7 g C/mi (9.1%) less CO<sub>2</sub>-C than does the PFI engine but 0.08 g C/mi more BC + OM, giving a ratio of CO<sub>2</sub>-C saved to BC + OM produced of 109:1. Figure 14 shows that, for a diesel vehicle to cool climate over 1 year, the ratio of CO<sub>2</sub>-C saved to BC + OM produced must exceed 5000–21,000:1, which is possible only if BC + OM emissions are reduced to 0.0004-0.0017 g/mi. For diesel to cool over 100 years, the ratio must exceed 220-500:1, which is possible only if BC + OM emissions are reduced to 0.02-0.04 g/mi. In sum, if the calculations here are correct, light-duty diesel cars meeting today's particulate standards and used daily will warm climate during the next 100+ years more than will gasoline cars.

[88] By 2005, E.U. particulate emission standards for light-duty vehicles will decrease to 0.025 g/km (0.04 g/mi). If all cars emitted at this limit, diesel would still warm climate more than would gasoline over a 100 year period (Figure 14). By 2004, California will implement Low Emission Vehicle (LEV) II standards for particulates of 0.01 g/mi (0.006 g/km) [the United States will implement a Tier II standard of up to 0.02 g/mi (0.012 g/km)]. If the tough California standards could be implemented worldwide, diesel would still warm climate more than would gasoline over a period of 13–54 years (Figure 14).

[89] In the United States and Europe, 99% of heavy-duty trucks and buses run on diesel. In the United States, <0.1% of light-duty vehicles run on diesel; in Europe, >25% run on diesel [Cohen and Nikula, 1999] (32.3% of new European cars in 2000 were diesel). Despite few U.S. diesel passenger vehicles, diesel consumption rates from all ground trans-

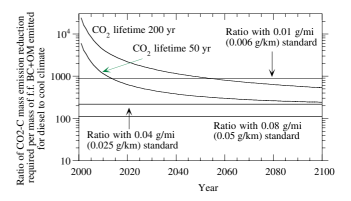


Figure 14. Ratio of CO<sub>2</sub>-C mass emission reduction required per mass of f.f. BC + OM emitted for diesel to cool climate. The curves were obtained by dividing the f.f. BC + OM temperature curve in Figure 1 by each CO<sub>2</sub> temperature curve [CO<sub>2</sub> (50 years) and CO<sub>2</sub> (200 years)] then multiplying the result by the current yearly emission rate of CO<sub>2</sub> (8100 Tg C/yr) and dividing by that of BC and associated OM from f.f. (5.1 Tg/yr BC + 10.1 Tg/yr OM). The figure shows that a yearly 1 Tg/yr decrease in f.f. BC + OM emissions will cool climate by 5000–21,000 times more than will a 1 Tg/yr decrease in CO<sub>2</sub>-C emissions during 1 year. After 100 years of continuous 1 Tg/yr decreases in both, the resulting ratio of f.f. BC + OM to CO<sub>2</sub>-C cooling is 220-500:1. Also shown (straight lines) are the ratios of CO<sub>2</sub>-C saved to f.f. BC + OM emitted for a modern diesel vehicle emitting 0.08, 0.04, and 0.01 g/mi BC + OM. The intersection of each of these straight lines with the two curves indicates the period of time during which diesel vehicles enhance global warming more than do gasoline vehicles under the given emission standard. In the case of the 0.08 and 0.04 g/mi standards, the period of time is >100 years.

portation sources (road, rail, inland waterways) in the United States are about 75-80% of those in Europe [International Energy Agency, 1999]. Table 4 shows that excise tax laws in all E.U. countries except the U.K. favor diesel, inadvertently promoting global warming. Some countries, (e.g., Finland, the Netherlands, Norway, and Sweden) also levy carbon taxes based on the carbon content of fuels. These taxes also favor diesel, since diesel releases less carbon per mile than does gasoline, but we calculate that the small release of BC + OM by diesel warms climate over 100 years than does the extra  $CO_2$  released by gasoline.

### 8. Conclusion

[90] Global model calculation in which 12 identifiable effects of aerosol particles on climate were accounted for were run. Results suggest that any emission reduction of f.f. BC + OM will slow global warming more than will any emission reduction of CO<sub>2</sub> or CH<sub>4</sub> for a specific period. When all f.f. BC + OM and anthropogenic CO<sub>2</sub> and CH<sub>4</sub> emissions are eliminated together, that period is 25–100 years. Historical net global warming can be attributed roughly to greenhouse gas plus f.f. BC + OM warming minus cooling due to other anthropogenic particles. Eliminating all f.f. BC + OM could eliminate 20–45% of such net warming (8–18% of total warming before cooling is

**Table 4.** Excise Taxes on Diesel and Gasoline in 1997 [European Commission, 1997]

	Excise tax on unleaded gasoline (ECU/1000 L)	Excise tax on diesel fuel (ECU/1000 L)	Ratio of tax on gasoline to tax on diesel
Belgium	510	292	1.75:1
Finland	536	310	1.73:1
Sweden	503	298	1.69:1
Netherlands	531	320	1.66:1
France	576	358	1.61:1
Germany	505	321	1.57:1
Denmark	447	308	1.45:1
Greece	363	252	1.44:1
Austria	416	291	1.43:1
Portugal	469	327	1.43:1
Spain	363	264	1.38:1
Italy	534	390	1.37:1
Luxembourg	349	254	1.37:1
Ireland	395	343	1.15:1
UK	501	501	1.00:1

subtracted out) within 3-5 years if no other change occurred. Reducing CO<sub>2</sub> emissions by a third would have the same effect, but after 50-200 years.

[91] Calculations also suggest diesel cars emitting continuously under the most recent U.S. and E.U. particulate standards may warm climate per distance driven over the next 100+ years more than do equivalent gasoline cars. If the estimates here are correct, fuel and carbon tax laws that favor diesel promote global warming. Toughening particulate emission standards to 0.006 g/km (0.01 g/mi), which is planned for California by 2004, does not change the conclusion, but it shortens the period over which diesel causes net warming to 13-54 years.

[92] Other practical strategies for reducing BC + OM are to phase out indoor biomass and coal burning and to improve particle collection from jet fuel and coal burning. Reducing BC + OM will not only slow global warming, but also improve human health. There may be no low threshold for health problems and mortality related to particles [Pope and Dockery, 1999], and about 2.7 million people die yearly from air pollution [WHO, 2000], many from indoor burning of biomass and coal and outdoor particles. In industrialized countries, the health costs of particulate matter are estimated to be \$200,000 to \$2.75 million per ton, making particles the most costly components of air pollution [Spadaro and Rabl, 2001]. Control of f.f. BC + OM alone will not eliminate long-term global warming. Elimination requires emission reductions in CO<sub>2</sub> and other greenhouse gases as well.

[93] Acknowledgments. This work was supported by the NASA New Investigator Program in Earth Sciences, the Environmental Protection Agency, the National Science Foundation, and the David and Lucile Packard Foundation and the Hewlett-Packard Company, I would also like to thank Don Wuebbles, Jim Hansen, Greg Carmichael, Matti Maricq, David Streets, Carey Jang, and John Bachmann for helpful comments and discussions

### References

Ackerman, A. S., B. Toon, D. E. Stevens, A. J. Heymsfield, V. Ramanathan, and E. J. Welton, Reduction of tropical cloudiness by soot, Science, 288, 1042-1047, 2000.

Ackerman, T. P., A model of the effect of aerosols on urban climates with

particular applications to the Los Angeles basin, J. Atmos. Sci., 34, 531-

Albrecht, B. A., Aerosols, cloud microphysics, and fractional cloudiness, Science, 245, 1227-1230, 1989

Allison, I., R. E. Brandt, and S. G. Warren, East Antarctic sea ice: Albedo, thickness distribution, and snow cover, J. Geophys. Res., 98, 12,417-

Andreae, M. O., and P. Merlet, Emission of trace gases and aerosols from biomass burning, Global Biogeochem. Cycles, 15(4), doi:10.1029/ 2000GB001382, 2002.

Andreae, M. O., T. W. Andreae, R. J. Ferek, and H. Raemdonck, Long range transport of soot carbon in the marine atmosphere, Sci. Total Environ., 36, 73-80, 1984.

Andres, R. J., and A. D. Kasgnoc, A time-averaged inventory of subaerial volcanic sulfur emissions, J. Geophys. Res., 103, 25,251-25,261, 1998. Angell, J. K., Comparison of surface and tropospheric temperature trends estimated from a 63-station radiosonde network, 1958-1998, Geophys. Res. Lett., 26, 2761-2764, 1999.

Akmaev, R. A., and V. I. Fomichev, A model estimate of cooling in the mesosphere and lower thermosphere due to the CO<sub>2</sub> increase over the last 3-4 decades, Geophys. Res. Lett., 27, 2113-2116, 2000

Arakawa, A., and V. R. Lamb, A potential enstrophy and energy conserving scheme for the shallow water equations, Mon. Weather Rev., 109, 18-36,

Beine, H., I. Allegrini, R. Sparapani, A. Ianniello, and F. Valentini, Three years of springtime trace gas and particle measurements at Ny-Alesund, Svalbard, Atmos. Environ., 35, 3645-3658, 2001.

Berresheim, H., M. O. Andreae, G. P. Ayers, R. W. Gillett, J. T. Merrill, V. J. Davis, and W. L. Chameides, Airborne measurements of dimethlylsulfide, sulfur dioxide, and aerosol ions over the Southern Ocean south of Australia, J. Atmos. Chem., 10, 341-370, 1990.

Bergstrom, R. W., and R. Viskanta, Modeling of the effects of gaseous and particulate pollutants in the urban atmosphere, part I, Thermal structure, J. Appl. Meteorol., 12, 901–912, 1973.

Berner, A., S. Sidla, Z. Galambos, C. Kruisz, R. Hitzenberger, H. M. ten Brink, and G. P. A. Kos, Modal character of atmospheric black carbon size distributions, J. Geophys. Res., 101, 19,559-19,565, 1996.

Bhugwant, C., H. Cachier, M. Bessafi, and J. Leveau, Impact of traffic on black carbon aerosol concentration at la Reunion Island (Southern Indian Ocean), Atmos. Environ., 34, 3463-3473, 2000.

Boccippio, D. J., K. L. Cummins, H. J. Christian, and S. J. Goodman, Combined satellite- and surface-based estimation of the intracloudcloud-to-ground lightning ratio over the continental United States, Mon. Weather Rev., 129, 108-122, 2001.

Bouwman, A. F., D. S. Lee, W. A. H. Asman, F. J. Dentener, K. W. van der Hoek, and J. G. J. Olivier, A global high-resolution emission inventory for ammonia, Global Biogeochem. Cycles, 11, 561-587, 1997

Brémond, M. P., H. Cachier, and P. Buat-Ménard, Particulate carbon in the

Paris region atmosphere, *Environ. Technol. Lett.*, 10, 339–346, 1989. Bretherton, C. S., E. Klinker, A. K. Betts, and J. Coakley, Comparison of ceilometer, satellite, and synoptic measurements of boundary layer cloudiness and the ECMWF diagnostic cloud parameterization scheme during ASTEX, J. Atmos. Sci., 52, 2736-2751, 1995.

Brorström-Lundén, E., A. Lindskog, and J. Mowrer, Concentrations and fluxes of organic compounds in the atmosphere of the Swedish west coast, Atmos. Environ., 28, 3605-3615, 1994.

Brunciak, P. A., J. Dachs, T. P. Franz, C. L. Gigliotti, E. D. Nelson, B. J. Turpin, and S. J. Eisenreich, Polychlorinated biphenyls and particulate organic/elemental carbon in the atmosphere of Chesapeake Bay, USA, Atmos. Environ., 35, 5663-5677, 2001.

Cachier, H., P. Buat-Ménard, M. Fontugne, and R. Chesselet, Long-range transport of continentally-derived particulate carbon in the marine atmosphere: Evidence from stable carbon isotope studies, Tellus, 38B, 161-177, 1986.

Cachier, H., M-P. Brémond, and P. Buat-Ménard, Determination of atmospheric soot carbon with a simple thermal method, Tellus, 41B, 379-390,

Cachier, H., M.-P. Brémond, and P. Buat-Ménard, Organic and black carbon aerosols over marine regions of the Northern Hemisphere, in Proceedings of the International Conference on Atmospheric Chemistry, edited by L. Newman et al., pp. 249-261, Brookhaven Natl. Lab., Upton, N. Y.,

Cachier, H., C. Liousse, A. Cachier, B. Ardouin, G. Polian, V. Kazan, and A. D. A. Hansen, Black carbon aerosols at the remote site of Amsterdam Island, in Fifth International Conference on Carbonaceous Aerosols, U.S. Dept. of Energy, Berkeley, Calif., 23-26 August 1994.

Chazette, P., and C. Liousse, A case study of optical and chemical ground apportionment for urban aerosols in Thessaloniki, Atmos. Environ., 35, 2497-2506, 2001.

Chowdhury, Z., L. S. Hughes, L. G. Salmon, and G. R. Cass, Atmospheric

- particle size and composition measurements to support light extinction calculations over the Indian Ocean, *J. Geophys. Res.*, 106, 28,597–28,605,2001
- Chylek, P., C. M. Banic, B. Johnson, P. A. Damiano, G. A. Isaac, W. R. Leaitch, P. S. K. Liu, F. S. Boudala, B. Winter, and D. Ngo, Black carbon: Atmospheric concentrations and cloud water content measurements over southern Nova Scotia, *J. Geophys. Res.*, 101, 29,105–29,110, 1996.
- Clairac, B., R. Delmas, B. Cros, H. Cachier, P. Buat-Menard, and J. Servant, Formation and chemical composition of atmospheric aerosols in an equatorial forest area, *J. Atmos. Chem.*, 6, 301–322, 1988.
- Clarke, A. D., Aerosol light absorption by soot in remote environments, Aerosol. Sci. Technol., 10, 161–171, 1989.
- Clarke, A. D., R. E. Weiss, and R. J. Charlson, Elemental carbon aerosols in the urban, rural, and remote-marine troposphere and in the stratosphere: Inferences from light absorption data and consequences regarding radiative transfer, *Sci. Total Environ.*, *36*, 97–102, 1984.
- Cohen, A. J., and K. Nikula, The health effects of diesel exhaust: Laboratory and epidemiologic studies, in *Air Pollution And Health*, edited by S. T. Holgate et al., pp. 707–745, Academic, San Diego, Calif., 1999.
- T. Holgate et al., pp. 707–745, Academic, San Diego, Calif., 1999. Cooke, W. F., and J. J. N. Wilson, A global black carbon aerosol model, *J. Geophys. Res.*, 101, 19,395–19,409, 1996.
- Cooke, W. F., C. Liousse, H. Cachier, and J. Feichter, Construction of a 1° × 1° fossil fuel emission data set for carbonaceous aerosol and implementation and radiative impact in the ECHAM4 model, *J. Geophys. Res.*, 104, 22,137–22,162, 1999.
- Del Delumyea, R., and A. Kalivretenos, Elemental carbon and lead content of fine particles from American and French cities of comparable size and industry, 1985, *Atmos. Environ.*, 21, 1643–1647, 1987.
- Derwent, R. G., D. B. Ryall, S. G. Jennings, T. G. Spain, and P. G. Simmonds, Black carbon aerosol and carbon monoxide in European regionally polluted air masses at Mace Head, Ireland during 1995–1998, Atmos. Environ., 35, 6371–6378, 2001.
- Dickerson, R. R., S. Kondragunta, G. Stenchikov, K. L. Civerolo, B. G. Doddridge, and B. N. Holben, The impact of aerosols on solar ultraviolet radiation and photochemical smog, *Science*, 278, 1997.
- Didyk, B. M., B. R. T. Simoneit, L. A. Pezoa, M. L. Riveros, and A. A. Flores, Urban aerosol particles of Santiago, Chile: Organic content and molecular characterization, *Atmos. Environ.*, 34, 1167–1179, 2000.
- Ding, P., and D. A. Randall, A cumulus parameterization with multiple cloud-base levels, J. Geophys. Res., 103, 11,341–11,353, 1998.
- Dzubay, T. G., R. K. Stevens, and P. L. Haagenson, Composition and origins of aerosol at a forested mountain in Soviet Georgia, *Environ. Sci. Technol.*, 18, 873–883, 1984.
- European Commission, Vehicle taxation in the European Union 1997, European Commission Directorate General XXI, background paper Ref XXI/ 306/98-EN, 1997.
- Ferek, R. J., J. S. Reid, P. V. Hobbs, D. R. Blake, and C. Liousse, Emission factors of hydrocarbons, halocarbons, trace gases and particles from biomass burning in Brazil, J. Geophys. Res., 103, 32,107–32,118, 1998.
- Food and Agricultural Organization (FAO), Soil Map of the World, Land and Water Dev. Div., FAO, Rome, Italy, 1995.
- Greenwald, T. J., G. L. Stephens, S. A. Christopher, and T. H. Vonder Haar, Observations of the global characteristics and regional radiative effects of marine cloud liquid water, *J. Clim.*, 8, 2928–2946, 1995.
- Gregory, G. L., et al., Air chemistry over the tropical forest of Guyana, J. Geophys. Res., 91, 8603-8612, 1986.
- Grenfell, T. C., S. G. Warren, and P. C. Mullen, Reflection of solar radiation by the Antarctic snow surface at ultraviolet, visible, and near-infrared wavelengths. *J. Geophys. Res.* 99, 18, 669–18, 684, 1994
- wavelengths, *J. Geophys. Res.*, 99, 18,669–18,684, 1994.
  Hack, J. J., J. T. Kiehl, and J. W. Hurrell, The hydrologic and thermodynamic characteristics of the NCAR CCM3, *J. Clim.*, 11, 1179–1206,
- Hamill, P., R. P. Turco, C. S. Kiang, O. B. Toon, and R. C. Whitten, An analysis of various nucleation mechanisms for sulfate particles in the stratosphere, *J. Aerosol. Sci.*, 13, 561–585, 1982.
- Hansen, J. E., and M. Sato, Trends of major climate forcing agents, *Proc. Natl. Acad. Sci.*, 93, 14,778–14,783, 2001.
- Hansen, A. D. A., B. A. Bodhaine, E. G. Dutton, and R. C. Schnell, Aerosol black carbon measurement at the South Pole: Initial results, 1986–1987, *Geophys. Res. Lett.*, 15, 1193–1196, 1988.
- Hansen, A. D. A., V. N. Kapustin, V. M. Kopeikin, D. A. Gillette, and B. A. Bodhaine, Optical absorption by aerosol black carbon and dust in a desert region of central Asia, *Atmos. Environ.*, 27A, 25,227–25,331, 1993
- Hansen, J., M. Sato, and R. Ruedy, Radiative forcing and climate response, J. Geophys. Res., 102, 6831–6864, 1997.
- Hansen, J., M. Sato, R. Ruedy, A. Lacis, and V. Oinas, Global warming in the twenty-first century: An alternative scenario, *Proc. Natl. Acad. Sci.*, 97, 9875–9880, 2000.

- Hayhoe, K., A. Jain, H. Pitcher, C. MacCracken, M. Gibbs, D. Wuebbles, R. Harvey, and D. Kruger, Costs of multigreenhouse gas reduction targets for the USA, *Science*, 286, 905–906, 1999.
- Haywood, J. M., D. L. Roberts, A. Slingo, J. M. Edwards, and K. P. Shine, General circulation model calculations of the direct radiative forcing by anthropogenic sulfate and fossil-fuel soot aerosol, *J. Clim.*, 10, 1562– 1577, 1997.
- He, K., F. Yang, Y. Ma, Q. Zhang, X. Yao, C. K. Chan, S. Cadle, T. Chan, and P. Mulawa, The characteristics of PM2.5 in Beijing, China, Atmos. Environ., 35, 4959–4970, 2001.
- Heidam, N. Z., P. Wahlin, and J. H. Christensen, Tropospheric gases and aerosols in Northeast Greenland, *J. Atmos. Sci.*, 56, 261–278, 1998.
- Heintzenberg, J., Size-segregated measurements of particulate elemental carbon and aerosol light absorption at remote Arctic locations, *Atmos. Environ.*, 16, 2461–2469, 1982.
- Heintzenberg, J., and A. Mészáros, Elemental carbon, sulfur, and metals in aerosol samples at a Hungarian regional air pollution station, *Idöjáràs*, 89, 313–319, 1985.
- Hillamo, R., V.-M. Kerminen, M. Aurela, T. Makela, W. Maenhaut, and C. Leck, Modal structure of chemical mass size distribution in the high Arctic aerosol, *J. Geophys. Res.*, 106, 27,555–27,571, 2001.
- Hitzenberger, R., and S. Tohno, Comparison of black carbon (BC) aerosols in two urban areas: Concentrations and size distributions, *Atmos. Envir*on., 35, 2153–2167, 2001.
- Hjellbrekke, A.-G. and J. E. Hanssen, Data report, 1996, 1, Annual summaries, *EMEP/CCC Rep. 1/98*, 85 pp., Norw. Inst. for Air Res., Lillestrom, 1998.
- Hoffert, M. I., A. J. Callegari, and C. T. Hsieh, The role of deep sea heat storage in the secular response of climate forcing, *J. Geophys. Res.*, 85, 6667–6679, 1980
- Im, J. S., V. K. Saxena, and B. N. Wenny, Temporal trends of black carbon concentrations and regional climate forcing in the southeastern United States, Atmos. Environ., 35, 3293–3302, 2001.
- Intergovernmental Panel on Climate Change (IPCC), *Third Assessment Report, Climate Change 2001: The Scientific Basis*, edited by J. T. Houghton et al., Cambridge Univ. Press, New York, 2001.
- International Energy Agency, Oil Information 1998, OECD/IED, Paris, 1999
- Jacobson, M. Z., Developing, coupling, and applying a gas, aerosol, transport, and radiation model to study urban and regional air pollution, Ph.D. Thesis, 436 pp., Univ. of Calif., Los Angeles, 1994.
- Jacobson, M. Z., Development and application of a new air pollution modeling system, part II, Aerosol module structure and design, Atmos. Environ., 31A, 131–144, 1997a.
- Jacobson, M. Z., Development and application of a new air pollution modeling system, part III, Aerosol-phase simulations, *Atmos. Environ.*, 31A, 587–608, 1997b.
- Jacobson, M. Z., Numerical techniques to solve condensational and dissolutional growth equations when growth is coupled to reversible reactions, Aerosol Sci. Technol., 27, 491–498, 1997c.
- Jacobson, M. Z., Improvement of SMVGEAR II on vector and scalar machines through absolute error tolerance control, *Atmos. Environ.*, 32, 791–796, 1998a.
- Jacobson, M. Z., Studying the effects of aerosols on vertical photolysis rate coefficient and temperature profiles over an urban airshed, *J. Geophys. Res.*, 103, 10,593–10,604, 1998b.
- Jacobson, M. Z., Fundamentals of Atmospheric Modeling, 656 pp., Cambridge Univ. Press, New York, 1999a.
- Jacobson, M. Z., Studying the effects of calcium and magnesium on sizedistributed nitrate and ammonium with EQUISOLV II, *Atmos. Environ.*, 33, 3635–3649, 1999b.
- Jacobson, M. Z., Isolating nitrated and aromatic aerosols and nitrated aromatic gases as sources of ultraviolet light absorption, *J. Geophys. Res.*, 104, 3527–3542, 1999c.
- Jacobson, M. Z., A physically-based treatment of elemental carbon optics: Implications for global direct forcing of aerosols, *Geophys. Res. Lett.*, 27, 217–220, 2000.
- Jacobson, M. Z., Strong radiative heating due to the mixing state of black carbon in atmospheric aerosols, *Nature*, 409, 695–697, 2001a.
- Jacobson, M. Z., Global direct radiative forcing due to multicomponent anthropogenic and natural aerosols, *J. Geophys. Res.*, 106, 1551–1568, 2001b.
- Jacobson, M. Z., GATOR-GCMM: A global through urban scale air pollution and weather forecast model, 1, Model design and treatment of subgrid soil, vegetation, roads, rooftops, water, sea ice, and snow, *J. Geophys. Res.*, 106, 5385-5402, 2001c.
- Jacobson, M. Z., GATOR-GCMM, 2, A study of day- and nighttime ozone layers aloft, ozone in national parks, and weather during the SARMAP Field Campaign, J. Geophys. Res., 106, 5403-5420, 2001d.
- Jacobson, M. Z., R. P. Turco, E. J. Jensen, and O. B. Toon, Modeling

- coagulation among particles of different composition and size, *Atmos. Environ.*, 28, 1327–1338, 1994.
- Janssen, N. A. H., D. F. M. Van Mansom, K. Van der Jagt, H. Harssema, and G. Hoek, Mass concentration and elemental composition of airborne particulate matter at street and background locations, *Atmos. Environ.*, 31, 1185–1193, 1997.
- Johansen, A. M., R. L. Siefert, and M. R. Hoffmann, Chemical composition of aerosols collected over the tropical North Atlantic Ocean, *J. Geophys. Res.*, 105, 15,277–15,312, 2000.
- Jones, P. D., D. E. Parker, T. J. Osborn, and K. R. Briffa, Global and hemispheric temperature anomalies: Land and marine instrumental records, in *Trends Online: A Compendium of Data on Global Change*, Carbon Dioxide Inf. Anal. Cent., Oak Ridge Natl. Lab., U.S. Dept. of Energy, Oak Ridge, Tenn., USA, 1999.
- Kadowaki, S., Characterization of carbonaceous aerosols in the Nagoya urban area, 1, Elemental and organic carbon concentrations and the origin of organic aerosols, *Environ. Sci. Technol.*, 24, 741–744, 1990.
- Kaneyasu, N., and S. Murayama, High concentrations of black carbon over middle latitudes in the North Pacific Ocean, *J. Geophys. Res.*, 105, 19,881–19,890, 2000.
- Keeler, G. J., S. M. Japar, W. W. Brachaczek, R. A. Gorse, J. M. Norbeck, and W. R. Pierson, The sources of aerosol elemental carbon at Allegheny Mountain, *Atmos. Environ.*, 24A, 2795–2805, 1990.
- Kerminen, V-M., K. Teinila, and R. Hillamo, Chemistry of sea-salt particles in the summer Antarctic atmosphere, *Atmos. Environ.*, 34, 2817–2825, 2000.
- Kettle, A. J., et al., A global database of sea surface dimethylsulfide (DMS) measurements and a procedure to predict sea surface DMS as a function of latitude, longitude, and month, *Global Biogeochem. Cycles*, 13, 399–444, 1999.
- Kim, B. M., S. Teffera, and M. D. Zeldin, Characterization of PM2.5 and PM10 in the South Coast Air Basin of Southern California, part 1, Spatial variations, J. Air Waste Manage. Assoc., 50, 2034–2044, 2000.
- Klein, S., Synoptic variability of low-cloud properties and meteorological parameters in the subtropical trade wind boundary layer, J. Clim., 10, 2018–2039, 1997.
- Knutson, T. R., T. L. Delworth, K. W. Dixon, and R. J. Stouffer, Model assessment of regional surface temperature trends (1949–1997), J. Geophys. Res., 104, 30,981–30,996, 1999.
- phys. Res., 104, 30,981–30,996, 1999.

  Kopeyken, V. M., V. N. Kapustin, and M. S. Pekur, Monitoring of soot aerosol in the atmosphere of Moscow, Izv. Acad. Sci. USSR, Atmos. Ocean. Phys. Engl. Transl., 29, 198–202, 1993.
- Krekov, G. M., Models of atmospheric aerosols, in *Aerosol Effects on Climate*, edited by S. G. Jennings, pp. 9–72, Univ. of Ariz. Press, Tucson, 1993.
- Krivacsy, Z., A. Hoffer, Zs. Sarvari, D. Temesi, U. Baltensperger, S. Nyeki, E. Weingartner, S. Kleefeld, and S. G. Jennings, Role of organic and black carbon in the chemical composition of atmospheric aerosol at European background sites, *Atmos. Environ.*, 35, 6231–6244, 2001.
- Light, B., H. Eicken, G. A. Maykut, and T. C. Grenfell, The effect of included particulates on the spectral albedo of sea ice, *J. Geophys. Res.*, 103, 27,739–27,752, 1998.
- Lighty, J. A., J. M. Veranth, and A. F. Sarofim, Combustion aerosols: Factors governing their size and composition and implications to human health, *J. Air Waste Manage. Assoc.*, 50, 1565–1618, 2000.
- Lin, J. J., and H.-S. Tai, Concentrations and distributions of carbonaceous species in ambient particles in Kaohsiung City, Taiwan, *Atmos. Environ.*, 35, 2627–2636, 2001.
- Liss, P. S., and L. Merlivat, Air-sea gas exchange rates, in *The Role of Air-Sea Exchange in Geochemical Cycling*, edited by P. Buat-Menard, pp. 113–127, 1986.
- Lobert, J. M., W. C. Keene, J. A. Logan, and R. Yevich, Global chlorine emissions from biomass burning: Reactive chlorine emissions inventory, *J. Geophys. Res.*, 104, 8373–8389, 1999.
- Lu, R., and R. P. Turco, Air pollutant transport in a coastal environment, part I, Two-dimensional simulations of sea-breeze and mountain effects, *J. Atmos. Sci.*, *51*, 2285–2308, 1994.
- Lyubovtseva, Y. S., and L. G. Yatskovich, Soot in the aerosols of different regions, J. Aerosol Sci., 20, 1269–1272, 1989.
- Maricq, M. M., R. E. Chase, D. H. Podsiadlik, and R. Vogt, Vehicle exhaust particle size distributions: A comparison of tailpipe and dilution tunnel measurements, SAE Technical Paper 1999-01-1461, SAE, 1999a.
- Maricq, M. M., D. H. Posiadlik, and R. C. Chase, Examination of the size-resolved and transient nature of motor vehicle particle emissions, *Environ. Sci. Technol.*, 33, 1618–1626, 1999b.
- Marland, G., T. A. Boden, R. J. Andres, A. L. Brenkert, and C. Johnston, Global, regional, and national CO<sub>2</sub> emissions, in *Trends Online: A Compendium of Data on Global Change*, Carbon Dioxide Inf. Anal. Cent., Oak Ridge Natl. Lab., U.S. Dept. of Energy, Oak Ridge, Tenn., USA, 1999

- Marticorena, B., G. Bergametti, B. Aumont, Y. Callot, and M. Legrand, Modeling the atmospheric dust cycle, 2, Simulation of Saharan dust sources, J. Geophys. Res., 102, 4387–4404, 1997.
- McNaughton, D. J., and R. J. Vet, Eulerian model evaluation field study (EMEFS): A summary of surface network measurements and data quality, *Atmos. Environ.*, 30, 227–238, 1996.
- Mellor, G. L., and T. Yamada, Development of a turbulence closure model for geophysical fluid problems, *Rev. Geophys. Space Phys.*, 20, 851–875, 1982.
- Mlawer, E. J., S. J. Taubman, P. D. Brown, M. J. Iacono, and S. A. Clough, Radiative transfer for inhomogeneous atmospheres: RRTM, a validated correlated-*k* model for the longwave, *J. Geophys. Res.*, *102*, 16,663–16,682, 1997.
- Modey, W. K., Y. Pang, N. L. Eatough, and D. J. Eatough, Fine particulate (PM2.5) composition in Atlanta, USA: Assessment of the particle concentrator-Brigham Young University organic sampling system, PC-BOSS, during the EPA supersite study, *Atmos. Environ.*, 35, 6493– 6502, 2001.
- Monahan, E. C., D. E. Spiel, and K. L. Davidson, A model of marine aerosol generation via whitecaps and wave disruption in oceanic whitecaps, in *Oceanic whitecaps and their role in air—sea exchange processes*, edited by E. C. Monahan and G. M. Niocaill, pp. 167–174, D. Reidel, Norwell, Mass., 1986.
- Mukai, H., Y. Ambe, K. Shibata, T. Muku, K. Takeshita, T. Fukuma, J. Takahashi, and S. Mizota, Long term variation of chemical composition of atmospheric aerosol on the Oki Islands in the Sea of Japan, *Atmos. Environ.*, 24A, 1379–1390, 1990.
- Nicholls, S., The dynamics of stratocumulus: Aircraft observations and comparisons with a mixed layer model, Q. J. R. Meteorol. Soc., 110, 783–820, 1984.
- Nijssen, B., R. Schnur, and D. P. Lettenmaier, Global retrospective estimation of soil moisture using the variable infiltration capacity land surface model, 1980–1993, *J. Clim.*, 14, 1790–1808, 2001.
- Noone, K. J., and A. D. Clarke, Soot scavenging measurements in Arctic snowfall, Atmos. Environ., 22, 2773–2778, 1988.
- O'Dowd, C. D., M. H. Smith, and S. G. Jennings, Submicron particle, radon, and soot carbon characteristics over the northeast Atlantic, *J. Geophys. Res.*, 98, 1123–1135, 1993.
- Offenberg, J. H., and J. E. Baker, Aerosol size distributions of elemental and organic carbon in urban and over-water atmospheres, *Atmos. Environ.*, 34, 1509–1517, 2000.
- Ohta, S., and T. Okita, Measurements of particulate carbon from the atmosphere, Sci. Total Environ., 18, 2439–2445, 1984.
- Olivier, J. G. J., A. F. Bouwman, C. W. M. Van der Maas, J. J. M. Berdowski, C. Veldt, J. P. J. Bloos, A. J. H. Visschedijk, P. Y. J. Zandveld, and J. L. Haverlag, Description of EDGAR Version 2.0: A set of global emission inventories of greenhouse gases and ozone-depleting substances for all anthropogenic and most natural sources on a per country basis on 1° × 1° grid, National Institute of Public Health and the Environment, RIVM report no. 771060 002/TNO-MEP report no. R96/119, 1996.
- Oltmans, S. J., and D. J. Hofmann, Increase in lower-stratospheric water vapour at a mid-latitude Northern Hemisphere site from 1981–1994, *Nature*, 374, 146–149, 1995.
- Park, S. S., Y. J. Kim, and K. Fung, Characteristics of PM2.5 carbonaceous aerosol in the Sihwa industrial area, South Korea, *Atmos. Environ.*, 35, 657–665, 2001.
- Parungo, F., C. Nagamoto, M-Y. Zhou, D. A. Hansen, and J. Harris, Aeolian transport of aerosol black carbon from China to the ocean, *Atmos. Environ.*, 28, 3251–3260, 1994.
- Polissar, A. V., Measurements of the soot mass concentration and particlesize distribution of atmospheric aerosol in the eastern Arctic, *Izv. Acad. Sci. USSR, Atmos. Ocean. Phys. Engl. Transl.*, 29, 66–73, 1993.
- Pope, C. A., III and D. W. Dockery, Epidemiology of particle effects, in *Air Pollution and Health*, edited by S. T. Holgate et al., pp. 673–705, Academic, San Diego, Calif., 1999.
- Pruppacher, H. R., and J. D. Klett, *Microphysics of Clouds and Precipitation*, 2nd rev. and enl. ed., Kluwer Acad., Norwell, Mass., 1997.
- Puxbaum, H., J. Rendl, R. Allabashi, L. Otter, and M. C. Scholes, Mass balance of the atmospheric aerosol in a South African subtropical savanna (Nylsvley, May 1997), *J. Geophys. Res.*, 105, 20,697–20,706, 2000.
- Quinn, P. K., D. J. Coffman, T. S. Bates, T. L. Miller, J. E. Johnson, K. Voss, E. J. Welton, and C. Neusus, Dominant aerosol chemical components and their contribution to extinction during the Aerosols99 cruise across the Atlantic, *J. Geophys. Res.*, 106, 20,783–20,809, 2001.
- Raunemaa, T., K. Kuuspalo, T. Alander, E. Tamm, A. Mirme, and V. Laine, Black carbon and aerosol in Böisto Island, J. Aerosol Sci., 24, S29–S30, 1993
- Reid, J. S., and P. V. Hobbs, Physical and optical properties of young smoke from individual biomass fires in Brazil, *J. Geophys. Res.*, 103, 32,013–32,030, 1998.

- Ruellan, S., and H. Cachier, Characterization of fresh particulate vehicular exhausts near a Paris high flow road, *Atmos. Environ.*, 35, 453–468, 2001.
- Savoie, D. L., J. M. Prospero, R. J. Larsen, F. Huang, M. Izaguirre,
  T. Huang, T. H. Snowdon, L. Custals, and C. G. Sanderson, Nitrogen
  and sulfur species in Antarctic aerosols at Mawson, Palmer Station, and
  March (King George Island), J. Atmos. Chem., 17, 95–122, 1993.
  Savoie, D. L., J. M. Prospero, R. Arimoto, and R. A. Duce, Non-sea-salt
- Savoie, D. L., J. M. Prospero, R. Arimoto, and R. A. Duce, Non-sea-salt sulfate and methanesulfonate at American Samoa, J. Geophys. Res., 99, 3587–3596, 1994.
- Schneider, T., and I. M. Held, Discriminants of twentieth-century changes in earth surface temperatures, J. Clim., 14, 249–254, 2001.
- Shauer, J. J., M. J. Kleeman, G. R. Cass, and B. R. T. Simoneit, Measurement of emissions from air pollution sources, 2, C<sub>1</sub> through C<sub>30</sub> compounds from medium duty trucks, *Environ. Sci. Technol.*, 33, 1578–1587, 1999.
- Smith, S. J., T. M. L. Wigley, and J. Edmonds, A new route toward limiting climate change, *Science*, *290*, 1109–1110, 2000.
- Spadaro, J. V., and A. Rabl, Damage costs due to automotive air pollution and the influence of street canyons, *Atmos. Environ.*, *35*, 4763–4775, 2001.
- Stelson, A. W., Urban aerosol refractive index prediction by partial molar refraction approach, *Environ. Sci. Technol.*, 24, 1676–1679, 1990.
- Stern, D. I., and R. K. Kaufman, Annual estimates of global anthropogenic methane emissions: 1860–1994, in *Trends Online: A Compendium of Data on Global Change*, Carbon Dioxide Inf. Anal. Cent., Oak Ridge Natl. Lab., U.S. Dept. of Energy, Oak Ridge, Tenn., USA, 1998.
- Strader, R., F. Lurmann, and S. N. Pandis, Evaluation of secondary organic aerosol formation in winter, *Atmos. Environ.*, 33, 4849–4863, 1999.
- Streets, D. G., S. Gupta, S. T. Waldhoff, M. Q. Wang, T. C. Bond, and B. Yiyun, Black carbon emissions in China, Atmos. Environ., 35, 4281– 4296, 2001
- Stott, P. A., S. F. B. Tett, G. S. Jones, M. R. Allen, J. F. B. Mitchell, and G. J. Jenkins, External control of 20th century temperature by natural and anthropogenic forcings, *Science*, 290, 2133–2137, 2000.
- Talbot, R. W., M. O. Andreae, H. Berresheim, P. Artaxo, M. Garstang, R. C. Harriss, K. M. Beecher, and S. M. Li, Aerosol chemistry during the wet season in central Amazonia: The influence of long-range transport, *J. Geophys. Res.*, 95, 16,955–16,969, 1990.
- Tang, I., Thermodynamic and optical properties of mixed-salt aerosols of atmospheric importance, *J. Geophys. Res.*, 102, 1883–1893, 1997.
   Temesi, D., A. Molnar, E. Meszaros, T. Feczko, A. Gelencser, G. Kiss, and
- Temesi, D., A. Molnar, E. Meszaros, T. Feczko, A. Gelencser, G. Kiss, and Z. Krivacsy, Size resolved chemical mass balance of aerosol particles over rural Hungary, *Atmos. Environ.*, 35, 4347–4355, 2001.
- Toon, O. B., and T. P. Ackerman, Algorithms for the calculation of scattering by stratified spheres, Appl. Opt., 20, 3657–3660, 1981.
- Toon, O. B., C. P. McKay, T. P. Ackerman, and K. Santhanam, Rapid calculation of radiative heating rates and photodissociation rates in inhomogeneous multiple scattering atmospheres, *J. Geophys. Res.*, 94, 16,287–16,301, 1989.
- Twomey, S. A., The effect of cloud scattering on the absorption of solar radiation by atmospheric dust, *J. Atmos. Sci.*, 29, 1156–1159, 1977.
- United States Geological Survey (USGS), U. Nebraska, Lincoln/European Commission's Joint Research Center 1-km resolution global landcover characteristics data base, derived from Advanced Very High Resolution Radiometer (AVHRR) data from the period April 1992 to March 1993, 1999
- Venkatram, A., and R. Viskanta, Radiative effects of elevated pollutant layers, J. Appl. Meteorol., 16, 1256–1272, 1977.

- Vogelmann, A. M., A. Robock, and R. G. Ellingson, Effects of dirty snow in nuclear winter simulations, J. Geophys. Res., 93, 5319–5332, 1988.
- Voldner, E. C., Y.-F. Li, M. T. Scholtz, and K. A. Davidson,  $1^{\circ} \times 1^{\circ}$  global SO<sub>x</sub> and NO<sub>x</sub> 2-level inventory resolved seasonally into emission sectors and point and area emissions sources, Global Emissions Inventory Activity (GEIA) website www.onesky.umich.edu/geia/, 1999.
- Walcek, C. J., and N. M. Aleksic, A simple but accurate mass conservative, peak-preserving, mixing ratio bounded advection algorithm with FOR-TRAN code, *Atmos. Environ.*, 32, 3863–3880, 1998.
- Walmsley, J. L., and M. L. Wesely, Modification of coded parameterizations of surface resistances to gaseous dry deposition, *Atmos. Environ.*, 30A, 1181–1188, 1996.
- Wang, M. Q., GREET 1.5: Transportation fuel-cycle model, volume 1, Methodology, development, use, and results, Argonne Natl. Lab. Rep. No. ANL/ESD-39, Vol. 1, 1999.
- Wang, C., and R. G. Prinn, On the roles of deep convective clouds in tropospheric chemistry, *J. Geophys. Res.*, 105, 22,269–22,297, 2000.
- Warren, S. G., Impurities in snow: Effects on albedo and snowmelt, *Ann. Glaciol.*, 5, 177-179, 1984.
- Warren, S. G., and A. D. Clarke, Soot in the atmosphere and snow surface of Antarctica, *J. Geophys. Res.*, 95, 1811–1816, 1990.
- Wesely, M. L., Parameterization of surface resistances to gaseous dry deposition in regional-scale numerical models, *Atmos. Environ.*, 23, 1293– 1304, 1989.
- Willison, M. J., A. G. Clarke, and E. M. Zeki, Seasonal variation in atmospheric aerosol concentration and composition at urban and rural sites in northern England, *Atmos. Environ.*, 19, 1081–1089, 1985.
- Wolff, G. T., M. S. Ruthkosky, D. P. Stroup, P. E. Korsog, M. A. Ferman, G. J. Wendel, and D. H. Stedman, Measurements of SOx, NOx and aerosol species on Bermuda, Atmos. Environ., 20, 1229–1239, 1986.
- World Health Organization (WHO), http://www.who, 2000.
- Wu, J., on of spume drops by the wind tearing of wave crests: The search for quantification, *J. Geophys. Res.*, 98, 18,221–18,227, 1993.
- Xie, P., and P. A. Arkin, Analyses of global monthly precipitation using gauge observations, satellite estimates, and numerical model predictions, *J. Clim.*, *9*, 840–858, 1996.
- Zappoli, S., et al., Inorganic, organic and macromolecular components of fine aerosol in different areas of Europe in relation to their water solubility, *Atmos. Environ.*, 33, 2733–2743, 1999.
  Zdunkowski, W. G., R. M. Welch, and J. Paegle, One dimensional numer-
- Zdunkowski, W. G., R. M. Welch, and J. Paegle, One dimensional numerical simulation of the effects of air pollution on the planetary boundary layer, *J. Atmos. Sci.*, *33*, 2399–2414, 1976.
- Zeng, X., R. E. Dickinson, A. Walker, M. Shaikh, R. S. DeFries, and J. Qi, Derivation and evaluation of global 1-km fractional vegetation cover data for land modeling, *J. Appl. Meteorol.*, 39, 826–839, 2000.
- Zhao, J., and R. P. Turco, Nucleation simulations in the wake of a jet aircraft in stratospheric flight, *J. Aerosol Sci.*, 26, 779–795, 1995.
- Zier, M., Mass concentrations of total dust and soot in the near surface air in east Germany, *J. Aerosol Sci.*, 22, S597–S600, 1991.

M. Z. Jacobson, Department of Civil and Environmental Engineering, Stanford University, Terman Engineering Center, Room M-13, Stanford, CA 94305-4020, USA. (jacobson@ce.stanford.edu)