Optical properties of boreal region biomass burning aerosols in central Alaska and seasonal variation of aerosol optical depth at an Arctic coastal site

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24	Earth's land surface or ~16 million square kilometers across northern Eurasia and North
25	America. The arctic and boreal zones currently store the globe's largest reservoir of soil
26	carbon, at 25-35% of the total, largely in organic soil layers in the permafrost [Bonan and
27	Shugart, 1989; Melillo et al. 1995]. Both climate change simulations of the effects of
28	increasing atmospheric greenhouse gas concentrations and recent observations show that
29	global temperature increases are the largest in the arctic and boreal regions [Soja et al.,
30	2007; IPCC, 2007]. As a result of these temperature increases (and drier conditions)
31	models predict future increases in area burned in the boreal zone. For example Flannigan
32	et al. [2005] simulated a 74-118% increase in burned area in Canada by 2100 in a 3XCO ₂
33	increase scenario. Kharuk et al. [2008] found a one-third reduction in fire return interval
34	time for larch dominated forests in Siberia from the 19 th to the 20 th century, related to
35	warming temperatures in northeast Siberia. Severe forest fires that also burn insulating
36	organic and peat soil layers result in deeper soil thawing [Kharuk et al., 2008] and
37	thereby may release large amounts of stored carbon into the atmosphere.
38	Concurrent with a significant warming trend, Kasischke and Turetsky [2006] found
39	that the annual boreal forest area burned in Alaska and Canada doubled from the
40	1960s/70s to the 80s/90s and the proportion of burning in the early and late growing
41	seasons increased. In Alaska seven of the eleven largest fires in a 56 year interval (1950-
42	2005) have burned since 1988, with the largest area burned on record occurring in 2004
43	and the third largest in 2005 [Soja et al., 2007]. Across the entire circumboreal zone the
44	frequency of extreme fire years has increased. Kasischke et al. [2002] have found that
45	high fire years in Alaska consist of larger fires occurring later in the growing season. In
46	low precipitation years peat burning is expected to increase as the summer advances due

strong boreal burning years they attribute ~80% of the global BC/snow forcing to be from
anthropogenic fossil fuel and biofuel sources.

72 Only recently have studies investigated the optical properties of boreal region biomass 73 burning aerosols in intensive burning years [Stohl et al., 2006b; Myhre et al., 2007]. 74 *Pfister et al.* [2008] discuss the need for better characterization of the optical properties 75 of boreal region biomass burning smoke particles, especially for the case of significant 76 peat burning. In this investigation we present an analysis of a long time series of aerosol 77 optical depth measurements from 1994 through 2008 at an AERONET sun-sky 78 radiometer site located in the boreal forest zone of central Alaska. In the extreme burning 79 years of 2004 and 2005 the AOD was very high, allowing for accurate characterization of 80 the spectral imaginary refractive index (absorption) and single scattering albedo (ω_0) 81 from almucantar retrievals. Additionally, we compare the particle size distributions and 82 ω_0 of these fine mode dominated smoke aerosols to the size distributions and absorption 83 of smoke aerosols from other major biomass burning regions. We also present AOD data 84 from a site on the Arctic Ocean coast in Alaska to compare the seasonality and frequency 85 of smoke transport to the Arctic (primarily in summer) to the springtime Arctic haze 86 impacts on optical depth. The Arctic haze phenomenon has been attributed primarily to 87 the long-distance transport of aerosols from industrial source regions (Shaw, 1995).

88

89 2. Instrumentation, Study Sites and Techniques

90 2.1 Study Region and Sites

91 The principal AERONET site analyzed in this study is the Bonanza Creek, Alaska site
92 located in the boreal forest biome of central Alaska (Figure 1). This is a National Science

116 every 15 minutes at 340, 380, 440, 500, 675, 870, 940, and 1020 nm (nominal 117 wavelengths). The direct sun measurements take ~ 8 seconds to scan all 8 wavelengths, 118 with a motor driven filter wheel positioning each filter in front of the detector. These 119 solar extinction measurements are then used to compute aerosol optical depth (AOD, τ_a) 120 at each wavelength except for the 940 nm channel, which is used to retrieve total 121 columnar (or precipitable) water vapor in centimeters. The filters utilized in these 122 instruments were ion assisted deposition interference filters with bandpass (full width at 123 half maximum) of 10 nm, except for the 340 and 380 nm channels at 2 nm. The estimated 124 uncertainty in computed τ_a , due primarily to calibration uncertainty, is ~0.010-0.021 for 125 field instruments (which is spectrally dependent with the higher errors in the UV; Eck et 126 al. [1999]). Schmid et al. [1999] compared τ_a values derived from 4 different solar 127 radiometers (including an AERONET sun-sky radiometer) operating simultaneously 128 together in a field experiment and found that the τ_a values from 380 to 1020 nm agreed to 129 within 0.015 (rms), which is similar to our estimated level of uncertainty in τ_a retrieval 130 for field instruments. The spectral aerosol optical depth data have been screened for 131 clouds following the methodology of *Smirnov et al.* [2000], which relies on the greater 132 temporal variance of cloud optical depth versus aerosol optical depth. The sky radiances 133 measured by the sun/sky radiometers are calibrated versus the 2-meter integrating sphere 134 at the NASA Goddard Space Flight Center, to an absolute accuracy of ~5% or better. 135 136

137 2.3 Inversion Methodology

to demonstrate successful retrievals of mode radii and the relative magnitude of modes
for various types of bimodal size distributions such as those dominated by a sub-micron
accumulation mode or distributions dominated by super-micron coarse mode aerosols. To
ensure sufficient sensitivity to aerosol absorption, only almucantar scans where
AOD(440nm)>0.4 [*Dubovik et al.*, 2000] were analyzed for the investigation of the
characteristics of spectral refractive indices and single scattering albedo.

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168 **3. Results and Discussion**

169 3.1 Temporal and Spectral Variability of AOD in central Alaska

170 3.1.1 Monthly and inter-annual variation in AOD and Angstrom exponent

171 The monthly climatology of 500 nm AOD and Angstrom Exponent (440-870 nm) at 172 Bonanza Creek, Alaska showing monthly means from multiple years of observations is 173 shown in Figure 2. The 440-870 nm Angstrom is computed from linear regression of ln 174 AOD versus ln λ scale at 440, 500, 670 and 870 nm. The multi-year monthly means are 175 computed as a mean of the individual monthly averages. The seasonal trend in monthly 176 average AOD shows a steady increase of AOD from near background levels in March 177 (~0.06) to values exceeding ~0.14 for June, July and August and then a rapid 2-month 178 decline down to background again (~0.05) in October. It should be noted that the inter-179 annual variability in AOD is extremely large (see Figure 3, discussed below) due to the 180 episodic nature of boreal fires with severe and numerous fires occurring in drought years, compared to few fires in wet years. Fine mode biomass burning aerosols dominate this 181 182 seasonality as the Angstrom exponent increases in summer (June-August) when most 183 burning occurs.

207 and 2008. However, in 2004 there were similar number of fires in July as August while 208 the AOD at Bonanza Creek was >2.5 times higher in August. Daily area burned estimates 209 for Alaska and Canada combined as shown by Stohl et al. [2006] reach a maximum from 210 late June through July, and much less area burned in August. This suggests the possibility 211 that factors other than area burned and numbers of fires are important in determining the 212 total atmospheric column AOD. Other factors in addition to area burned that would 213 influence the AOD are meteorological conditions (including wind speed and direction, 214 atmospheric stability, and precipitation), intensity of burning since intense fires enhance 215 convection that may loft smoke to higher altitudes where winds are higher, and the types 216 of fuel burned and the phase of combustion [*Reid et al.*, 2005b]. It is noted that AOD 217 measurements at a point location (Bonanza Creek) are also strongly affected by the 218 location of the fires in the entire state of Alaska relative to the wind direction and other 219 meteorological factors. Satellite hot spot remote sensing robustly detects fires in the high 220 temperature flaming phase of combustion but it is much more difficult to detect lower 221 temperature smoldering phase fires. However the flaming phase is typically of relatively 222 short duration, while the larger diameter woody fuels in a forest and the organic soils and 223 peatlands may burn for several days in the smoldering phase. As previously mentioned, 224 Turquety et al. [2007] estimated that the proportion of carbon monoxide emissions was 225 double the proportion of peat area burned (~37% of the carbon monoxide emissions from 226 only 17% of the area burned) in 2004 for fires in Alaska and Canada. This suggests that 227 peat burning may possibly also have resulted in a disproportionate amount of total 228 aerosol emissions, relative to the total area burned.

variation in source strength and due to the transport, stagnation, and removal effects ofregional meteorology.

253 Based on the assumption that aerosol size distributions are bimodal, O'Neill et al. 254 (2001, 2003) have developed a spectral deconvolution algorithm (SDA) that utilizes 255 spectral total extinction AOD data to infer the component fine and coarse mode optical 256 depths. An additional fundamental assumption of the algorithm is that the coarse mode 257 Angstrom exponent and its derivative are both close to zero. The Angstrom exponent α 258 and the spectral variation of α (as parameterized by $\alpha' = d\alpha/d\ln\lambda$) are the measurement 259 inputs to the SDA. These continuous-function derivatives (usually computed at a reference wavelength of 500 nm) are derived from a second order fit of ln τ_a versus ln λ 260 261 (Eck et al., 1999). The spectral AODs employed as input to the SDA were limited to the 262 six CIMEL wavelengths ranging from 380 to 1020 nm. Figure 6 shows the time series of 263 fine and coarse mode daily average AOD at 500 nm from the SDA algorithm for June 1 264 through September 12, 2004 at Bonanza Creek. These are level 2 cloud screened data that 265 have had the SDA algorithm applied to the AOD spectra. It is noted that the coarse mode 266 is typically very low and nearly constant, while the fine mode AOD from biomass 267 burning exhibits very large day-to-day variability. Only 2 days show the coarse mode 268 AOD to be significantly higher than the fine mode, possibly due to dust transported from 269 Asia or from residual cloud contamination. We assume that the fine mode aerosols at 270 Bonanza Creek are dominated by biomass burning smoke since there are numerous fire 271 hot spots observed (Figure 1), combined with a lack of any other significant regional 272sources of fine mode particles.

295	The SDA algorithm computed daily average fine mode fractions (FMF) for the
296	Bonanza Creek site are plotted versus $\alpha_{380-500}$ in Figure 8, and compared with
297	measurements made in Brazil and Zambia (from the same observations as shown in
298	Figure 7). For the same range of FMF, $0.8 - 1.0$, the $\alpha_{380-500}$ is typically significantly
299	lower for the Alaska site than the Brazil and Zambia biomass burning sites, again
300	suggesting much larger accumulation mode particles in Alaska. In addition to the greater
301	percentage of smoldering combustion for the forest regions with woody fuels (Alaska and
302	Brazil) other possible reasons for larger accumulation mode particles include higher
303	AOD levels (Figure 7) in Alaska and Brazil that increase the coagulation rate since
304	aerosol concentrations are higher. Another possible factor in creating large accumulation
305	particles might be the characteristics of smoke from peat burning in Alaska [Reid et al.,
306	2005a], since extensive areas of peat lands (in addition for forests) were observed to burn
307	during the summer of 2004.
308	
309	3.2 Retrievals of aerosol size distribution and single scattering albedo in central
310	Alaska
311	3.2.1 Volume size distributions
312	The almucantar retrievals of aerosol volume size distributions for the Bonanza Creek
313	site in 2004 and 2005 for scans where AOD(440nm)>0.4 are shown in Figure 9a. These
314	averages are plotted as a function of AOD(440nm) with AOD bins from 0.4-0.8, >0.8-
315	1.0,,>2.4-2.8, >2.8 resulting in averages of from 9 to 34 almucantars per bin. At all
316	AOD levels the retrievals show the dominance of fine mode aerosols, and the $\alpha_{440-870}$ for
317	the bins range from 1.74 at the lowest AOD(440) average of 0.53 to 1.36 at the highest

341	exhibiting $r_v \sim 0.21 \ \mu m$ and transported smoke from a Russian peat/forest fire showing
342	retrieved r_{ν} of 0.28 μm (Version 2 retrievals as compared to somewhat smaller radius
343	values reported from Version 1 in Eck et al. [2003a]). These however were aged aerosol
344	events with smoke age > 2 days due to long distance transport from distinct source
345	regions. However, AERONET measurements made in Moscow on September 7, 2002
346	$(AOD(440) \sim 2.5)$ near to a fire that was predominately burning peat had large
347	accumulation mode particles of ~0.22 μ m radius despite relatively short transport
348	distance and therefore likely little aging. Therefore, the large radius of the smoke aerosol
349	measured in late summer in Alaska in 2004 and 2005 possibly resulted in part from the
350	smoldering combustion of peat fuels in addition to the high aerosol concentrations that
351	would result in greater coagulation, condensation and secondary production rates.
352	In comparison to the Alaska smoke we show the aerosol volume size distribution
353	retrievals for biomass burning aerosols from the Mongu, Zambia AERONET site in
354	Figure 11a. These retrievals are from September (peak burning month) data only for the
355	years 1997-2005, and shown for two AOD levels, ~0.5 and ~1.3 at 440 nm. Both the
356	mean sizes and the shift of fine mode radius as AOD increases in Zambia are relatively
357	small (radius ~0.14 to 0.16 μ m) as compared to the boreal smoke measured in Alaska.
358	Smoke in Brazil (southern Amazonia; not shown) exhibited slightly larger fine mode
359	radius (~0.15 to 0.17 µm; Schafer et al., 2008) at these AOD levels than Zambia, possibly
360	due to more smoldering combustion of woody fuels and higher aerosol concentrations
361	that may have lead to greater coagulation rates. In addition to the relatively small change
362	in volume median radius in Mongu, Zambia, the width of the fine mode size distribution
363	is narrower for the Zambia smoke than for the Alaska smoke. The geometric standard

.

386 The average single scattering albedo in 2004 and 2005 at Bonanza Creek is high 387 (weak absorption) for biomass burning aerosols, with most values ranging from ~0.96 to 388 0.97 (Figure 9b). Only four of the individual retrievals out of a total of 124 almucantars 389 with AOD(440nm)>0.4 in 2004 and 2005 had ω_0 less than 0.935 at 440 nm, with the 390 lowest at 0.909. The highest retrieved value of ω_0 at 440 nm was 0.996. For all but the 391 lowest AOD level the average ω_0 at 440 nm is slightly lower (by ~0.01) than that at 675 392 nm, and the ω_0 from 675 nm through 1020 nm are relatively constant (Figure 9b). This 393 spectral dependence of ω_0 appears to be anomalous for biomass burning aerosols, as the 394 single scattering albedo typically decreases with increasing wavelength in both 395 measurements and retrievals [Reid et al., 2005b]. For example, Dubovik et al. [2002] 396 show this typical wavelength dependence for biomass burning aerosols from four major 397 regions: Amazonian forest, S. American cerrado (savanna-like), African savanna in 398 Zambia, and boreal forest (primarily Canada). However, the boreal forest data set in 399 Dubovik et al. [2002] does not include events with as high AOD as occurred in Alaska in 400 2004 and 2005, nor does it include observations with significant peat burning. For all 401 other years from 1994-2007 (excluding 2004 and 2005) there were a total of only 14 402 almucantar retrievals at Bonanza Creek with AOD(440nm)>0.4. Two of these had ω_0 at 403 400 nm of ~0.86 which suggests flaming phase crown fires, while the other 12 had values 404 ranging from 0.92 to 0.98. The mean of these 14 almucantars was 0.94 at 440 nm and 405 within less than 0.005 of the values given by *Dubovik et al.* [2002] at all 4 wavelengths 406 (Dubovik's values were a mean for boreal forest biomass burning aerosols). Similar wavelength dependence of ω_0 to the Alaskan smoke of 2004 and 2005 was 407 408 observed however for the previous mentioned case of peat burning smoke in Moscow on

432 higher and relatively constant (~0.72). For the burning of lignite fuel Bond et al. [1999] 433 also measured greater fine particle absorption at shorter wavelengths (450 nm), implying 434 larger imaginary refractive index at shorter wavelengths, possibly due to absorption by 435 organic carbon. It is noted that their laboratory combustion of this lignite fuel occurred 436 partially in the smoldering phase. However it is noted that the absorption properties of 437 organic carbon are not well known and therefore are a topic of much recent research. 438 Another factor that contributes to the relatively constant spectral single scattering 439 albedo of the Alaskan smoke is the much larger size and wider distribution of the fine 440 mode particle radius, as compared to particles from most other biomass burning regions. 441 Because the scattering cross section (or hence scattering optical depth) increases more 442 rapidly than absorption cross section with increasing particle size the SSA also increases 443 (in the absence of variations in refractive index) with increasing particle size. This trend 444 is the optical equivalent of stating (as above) that SSA typically decreases with increasing 445 wavelength. However this trend is less extreme at larger particle sizes because the SSA 446 approaches unity at a lesser rate with respect to increasing particle size (or decreasing 447 wavelength).

In situ measurements from nephelometer and particle soot absorption photometer data at the surface in Barrow, Alaska on July 3-4, 2004 yielded an aerosol single scattering albedo of 0.96 at 550 nm [*Stohl et al.*, 2006; *Stone et al.*, 2008], for a case of very high smoke AOD advected from fires in central Alaska and the Yukon. This is essentially equal to the mean AERONET retrievals of ω_0 interpolated to 550 nm for the total column aerosol in central Alaska at Bonanza Creek (Figure 9a). For agricultural smoke originating in Europe and subsequently advected to Svalbard in the Arctic, *Myhre et al.*

478 conditions, *Lewis et al.* [2008] measured AAE as high as 2.5 (532 to 870 nm) at high ω_0 479 (near unity) while AAE values approached 1.0 for $\omega_0 < 0.8$ at 532nm. The measured 480 organic carbon fraction to total carbon was highest for the smoke with the highest AAE 481 and ω_0 , thereby suggesting that the enhanced absorption AAE results from light 482 absorbing organic carbon.

483

484 **3.3 Seasonal variation of AOD in the coastal Arctic at Barrow, Alaska**

In this section we present data from the Arctic AERONET site located at Barrow, 485 486 Alaska on the Beaufort Sea coast (Figure 1). The data collection at this site has many 487 more gaps than for the Bonanza Creek site in central Alaska due to its more severe 488 weather, which sometimes resulted in instrument electronic or mechanical problems. This 489 occurred especially in earlier years before a modified version of the CIMEL that 490 incorporated heating elements was deployed. Additionally there is more persistent cloud 491 cover at this site than at Bonanza Creek, resulting in fewer observations of the sun, 492 therefore less AOD measurements and very few almucantar scans of sky radiance 493 distribution. Data were acquired in 1999, 2002, 2004, 2005, 2006, 2007, and 2008 494 however measurements were only made during the peak Arctic haze month of April in 495 three years, 2002, 2005 and 2008, when data collection began in late March or early 496 April. All years had monitoring from July through September, and four of the six years 497 had data from May through September, thus covering the biomass burning season. 498 Therefore due to the numerous gaps in data acquisition, the AERONET data presented 499 here for Barrow cannot be considered a fully representative monitoring record, especially 500 for the spring arctic haze season. These data gaps therefore preclude any analysis of

524	solar zenith angle. Stohl et al. [2006a] have shown from transport modeling that this
525	smoke event originated from the fires located in central Alaska and the Canadian Yukon,
526	and that the smoke continued to be transported beyond Barrow and deep into the arctic.
527	Their simulations suggested that smoke from this event reached the North Pole on July 8,
528	2004 although cloud cover precluded verification from satellite images. The AERONET
529	site located at Resolute Bay, Canada (74 ⁰ 44' N, 94 ⁰ 54' W; ~1950 km ENE from
530	Barrow) measured smoke AOD on July 5, 2004 as high as 2.3 at 500 nm ($\alpha_{440-870} > 1.3$)
531	from this same arctic transport event. A Terra MODIS image (Figure 7 in Stohl et al.
532	(2006)) shows widespread smoke from Alaska and the Yukon through the Arctic islands
533	of Canada on July 5, 2004.
534	Figure 12b is the same as 12a but with the extreme AOD event of July 3, 2004
535	excluded. Individual daily averages of AOD are shown as well as 20 day interval means
536	(means computed with the one extreme day removed also). Seasonality of AOD is
537	evident, with the highest 20-day averages of AOD occurring during the arctic haze season
538	from late March through mid-May. These higher average AOD values result partly from
539	the lack of measured low background AOD in the spring, when values are rarely lower
540	than 0.07. In contrast the daily average AOD during summer months is often <0.04 and
541	as low as 0.02. However daily average AOD on some summer days, as high as or higher
542	than the spring Arctic haze AOD, from biomass burning aerosols result in mean values of
543	AOD during the summer that are elevated significantly above background levels. Again it
544	is emphasized that due to a much less extensive data record at Barrow (as compared to
545	Bonanza Creek), and since the extreme Alaska burning years of 2004 and 2005 are

monthly average Angstrom Exponent (440-870 nm) at both sites, 1.25 at Barrow and 1.10
at Bonanza Creek.

571 Shaw [1982] measured AOD from sunphotometer at Barrow during the mid to late 1970s and computed a March-April mean of 0.135 at 500 nm. Therefore the mean AOD 572 573 measured by AERONET for the spring arctic haze (and smoke) at Barrow for 2002, 2005 574 and 2008 were somewhat higher than that measured ~25 years earlier. Bodhaine and 575 Dutton [1993] presented measurements of AOD at Barrow computed from broadband 576 (300-690 nm) pyrheliometer measurements for the years 1977-1992, with estimates of 577 volcanic aerosol optical depth removed. They show relatively low AOD in 1980 and 578 1981 and a significant downward trend from 1982 (peak year) to 1992 that they suggest 579 may have resulted from the reduction of emissions in the Soviet Union and Europe during 580 that era. Continued monitoring of AOD at Barrow is important for understanding arctic 581 haze magnitude and trends. Ground based photometric measurements of AOD at 532 nm 582 at the arctic island of Spitsbergen (~79N, 12E) by Herber et al. [2002] from 1991 583 through 1999 showed a gradual increase in AOD of $\sim 9\%$ over the 9 year interval. The 584 relative sparseness of long-term records of AOD at Arctic locations coupled with the 585 possibility of different regional influences (sources and meteorology) make it very 586 difficult to assess trends of aerosol loading across the entire arctic region.

587

588 4. Summary and Conclusions

589

Aerosol optical properties data acquired from monitoring at two AERONET sites inAlaska were investigated. Data from long-term monitoring at a central Alaska boreal

614	3.	Absorption by the smoke aerosol in Bonanza Creek in 2004 and 2005 was very
615		weak, with retrieved single scattering albedo ranging from ~0.96 to 0.97, along
616		with relatively flat spectral dependence. These high single scattering albedos
617		result from small values of the imaginary index of refraction, implying low black
618		carbon fraction probably due to predominately smoldering combustion, coupled
619		with large fine mode particle radius which results in greater scattering efficiency
620		(increased ω_o amplitude) and reduced wavelength dependence of the ω_o .
621		Additionally, the single scattering albedo at 440 nm was ~0.01 lower than at the
622		longer wavelengths due to a somewhat larger imaginary refractive index at 440
623		nm, which is possibly due to enhanced short wavelength absorption by organic
624		carbon aerosols. This also suggests the possibility that smoke from peat burning
625		(smoldering combustion) had a significant influence on aerosol emissions.
626	4.	Although AERONET monitoring at the Arctic coastal site of Barrow from 1999 to
627		2008 was often interrupted and not complete enough to be considered a
628		representative climatology, some seasonal characteristics of AOD were
629		nonetheless evident. The average AOD in the spring (late March through late
630		May) is higher than the average AOD in the summer. Even though several
631		individual daily mean values in summer are significantly higher (from transported
632		biomass burning smoke) than most daily means in spring, the lack of very low
633		background AOD levels in spring, due primarily to persistent industrial arctic
634		haze, resulted in higher mean AOD in spring.
635		

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Figure 5. Remotely sensed fire counts for the Alaska region from the MODIS sensor onthe Terra satellite for the years 2002 through 2008.

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Figure 6. Spectral Deconvolution Algorithm (SDA) computed daily average fine and

coarse mode AOD versus day of the year from June 1, 2004 through September 12, 2004

846 at Bonanza Creek, Alaska.

847

Figure 7. A comparison of the daily average 380-500 nm Angstrom exponents as a

function of 500 nm AOD for the Bonanza Creek data from 2004 versus the AERONET

data from the major tropical region biomass burning sites of ABRACOS Hill, Brazil

851 (2002 data; southern Amazonia) and Mongu, Zambia (2004; southern Africa savanna

burning region). For all three sites the data for the June through October biomass burningseasons are shown.

854

855 Figure 8. Spectral Deconvolution Algorithm (SDA) computed daily average fine mode

856 fraction (FMF) for the Bonanza Creek site plotted versus $\alpha_{380-500}$ and compared with

857 measurements made in Brazil and Zambia (from the same observations as shown in

858 Figure 6).

859

Figure 9. Almucantar retrievals of a) aerosol volume size distributions b) single scattering albedo, and c) imaginary part of the refractive index from the Bonanza Creek site in 2004 and 2005 for scans where AOD(440nm)>0.4. These averages are plotted as a function of













Figure 6. Spectral Deconvolution Algorithm (SDA) computed daily average fine and
coarse mode AOD versus day of the year from June 1, 2004 through September 12, 2004
at Bonanza Creek, Alaska.





albedo, and c) imaginary part of the refractive index from the Bonanza Creek site in 2004
 and 2005 for scans where AOD(440nm)>0.4. These averages are plotted as a function of

1016 AOD(440nm) for AOD bins from 0.4-0.8, >0.8-1.0,...,>2.4-2.8, >2.8 resulting in

1017 averages of from 9 to 35 almucantar scans per bin.







1072Day of Year1073Figure 12. a) The daily average 500 nm AOD measured at the Barrow AERONET site as1074a function of the day of the year for all monitoring during the 1999 through 2008 time1075interval. b) The same as in a) but with the single outlier point of AOD (500 nm) of 3.41076from July 3, 2004 removed, and with 20 day averages computed.