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# Improved estimate of the policy-relevant background ozone in the United States using the GEOS-Chem global model with 1/2° × 2/3° horizontal resolution over North America

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- 1 Improved estimate of the policy-relevant background ozone in the United States
- 2 using the GEOS-Chem global model with  $1/2^{\circ} \times 2/3^{\circ}$  horizontal resolution over
- 3 North America
- 4
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- 24

### 24 Abstract

25

26 The policy-relevant background (PRB) ozone is defined by the US Environmental 27 Protection Agency (EPA) as the surface ozone concentration that would be present over 28 the US in the absence of North American anthropogenic emissions. It is intended to 29 provide a baseline for risk and exposure assessments used in setting the National 30 Ambient Air Quality Standard (NAAQS). We present here three-year statistics (2006-31 2008) of PRB ozone over the US calculated using the GEOS-Chem global 3-D model of 32 atmospheric composition with  $1/2^{\circ} \times 2/3^{\circ}$  horizontal resolution over North America and 33 adjacent oceans  $(2^{\circ} \times 2.5^{\circ})$  for the rest of the world). We also provide estimates of the US 34 background (no anthropogenic US emissions) and natural background (no anthropogenic 35 emissions worldwide and preindustrial methane). PRB is particularly high in the 36 intermountain West due to high elevation, arid terrain, and large-scale subsidence. We 37 present for this region a detailed model evaluation showing that the model is successful in 38 reproducing ozone exceedances up to 70 ppby. However, the model cannot reproduce 39 PRB-relevant exceptional events associated with wildfires or stratospheric intrusions. The 40 mean PRB estimates for spring-summer are  $27 \pm 8$  ppby at low-altitude sites and  $40 \pm 7$ 41 ppby at high-altitude sites. These include a mean enhancement from intercontinental 42 pollution and anthropogenic methane of 9 ppby at low-altitude sites and 13 ppby at high-43 altitude sites. The PRB is higher than average when ozone exceeds 60 ppby, particularly in the intermountain West. The annual 4<sup>th</sup>-highest PRB values in the intermountain West 44 45 are typically 50-60 ppby, as compared to 35-45 ppby in the East or on the West Coast. 46 Our PRB estimates are on average 4 ppbv higher than in previous GEOS-Chem studies 47 and this may reflect higher lightning, increasing Asian emissions, and improved model 48 resolution. 49

50 **Keywords:** Ozone; background ozone; policy relevant background; air quality standard 51

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### 53 1. Introduction

54

55 The US Environmental Protection Agency (US EPA, 2006) defines the policy-relevant-56 background (PRB) for ozone air quality as the surface ozone concentration that would be 57 present in the US in the absence of anthropogenic emissions from North America 58 (defined as the ensemble of the US, Canada, and Mexico). The PRB is used in the setting 59 of the National Ambient Air Quality Standard (NAAQS) to estimate the maximum ozone 60 reduction that could be achieved through North American emission controls. It provides 61 a baseline for risk and exposure assessments. The present US NAAQS is 75 ppbv (annual 62 4<sup>th</sup>-highest daily maximum 8-h average concentration), but the EPA is considering 63 decreasing it to a value in the 60-70 ppbv range. As the standard becomes more stringent 64 and approaches the PRB, accurate specification of the PRB becomes increasingly 65 important

66

67 Ozone is produced in the troposphere by photochemical oxidation of CO and volatile

- organic compounds (VOCs) in the presence of nitrogen oxides (NO<sub>x</sub>  $\equiv$  NO + NO<sub>2</sub>). It is
- also transported to the troposphere from the stratosphere. The NO<sub>x</sub>, CO, and VOC

70 precursors of ozone have major anthropogenic sources from fuel combustion as well as 71 natural sources including lightning, wildfires, and the biosphere. Ozone has a lifetime of 72 only a few days in the continental boundary layer but weeks in the free troposphere (Y. 73 Wang et al., 1998; Fiore et al., 2002). Ozonesonde, aircraft, and satellite observations 74 show typical ozone concentrations of 50-70 ppby in the free troposphere over North 75 America (Thompson et al., 2007; L. Zhang et al., 2010), with frequent occurrence over 80 76 ppby in plumes from intercontinental pollution, fires, and stratospheric intrusions (Heald 77 et al., 2003; Nowak et al., 2004; Bertschi and Jaffe, 2005; Liang et al., 2007; Thompson 78 et al., 2007; Oltmans et al., 2010). Subsidence of this high-ozone air to the surface could 79 result in PRB values approaching or exceeding the NAAQS (Jaffe, 2011). However, 80 ozone decreases during entrainment into the boundary layer because of dilution,

- 81 deposition, and chemical loss (Fiore et al., 2002; Hudman et al., 2004; L. Zhang et al., 2009; Cooper et al., 2011).
- 82
- 83

84 A region of particular interest from a PRB perspective is the intermountain West.

85 extending between the Sierra Nevada/Cascades Mountains to the west and the Rocky

86 Mountains to the east. This region features elevated plateaus and mountains with surface 87 elevations typically in excess of 1.5 km, arid terrain, and large-scale subsidence. As a

88 result, background ozone there is higher than in the eastern US (Lefohn et al., 2001; Fiore

89 et al., 2002; Jaffe, 2011). Exceptionally high ozone events have been observed in

90 association with stratospheric intrusions (Langford et al., 2009). Positive correlations

91 have been observed between ozone and regional wildfires (Jaffe et al., 2008; Jaffe, 2011).

92 There is evidence that ozone inflow from the Pacific to the western US has been

93 increasing over the past decades (Lin et al., 2000; Jaffe and Ray, 2007; Parrish et al.,

94 2009; Cooper et al., 2010). This could reflect increasing Siberian wildfires (Jaffe et al.,

- 95 2004) and Asian pollution (L. Zhang et al., 2008; Cooper et al., 2010).
- 96

97 The PRB ozone is not an observable quantity, if only because of the contribution of North 98 American anthropogenic sources to the northern mid-latitudes ozone background. It 99 needs to be calculated with a global model of atmospheric composition that is evaluated 100 with observations at remote sites where the PRB drives much of the variability. Fiore et 101 al. (2003) previously used the GEOS-Chem CTM with  $2^{\circ} \times 2.5^{\circ}$  global horizontal 102 resolution to estimate PRB ozone over the US. H. Wang et al. (2009) updated those 103 estimates by using a  $1^{\circ} \times 1^{\circ}$  nested continental-scale version of GEOS-Chem, and also 104 estimated the US background ozone (defined by zeroing anthropogenic US emissions). Here we present a further update of PRB, US background, and natural background ozone 105 106 estimates with a 3-year (2006-2008) GEOS-Chem simulation at  $1/2^{\circ} \times 2/3^{\circ}$  resolution 107 featuring a number of improvements over previous versions. Our motivation for this 108 work is to assist the EPA in its revision of the ozone NAAOS, scheduled to be released in 2014. We include a detailed model evaluation in the intermountain West where elevated 109 PRB is of particular relevance to the NAAQS.

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- 111

#### 112 2. Model description

113

114 We use the GEOS-Chem 3-D global model of atmospheric composition (v8-02-03; 115 http://acmg.seas.harvard.edu/geos/). The model is driven by GEOS-5 assimilated

- 116 meteorological fields from the NASA Global Modeling and Assimilation Office
- 117 (GMAO). Meteorological fields in the GEOS-5 data have a temporal resolution of 6
- 118 hours (3 hours for surface variables and mixing depths) and a horizontal resolution of
- 119 1/2° latitude by 2/3° longitude. GEOS-Chem includes a detailed simulation of
- 120 tropospheric ozone-NOx-VOC-aerosol chemistry. The ozone simulation over the US and
- 121 adjacent oceans has been previously evaluated with measurements from surface sites
- 122 (Fiore et al., 2002, 2003; Goldstein et al., 2004; H. Wang et al., 2009), aircraft (Hudman
- 123 et al., 2007; L. Zhang et al., 2008; Walker et al., 2010), ozonesondes (Li et al., 2002,
- 124 2005), and satellites (Parrington et al., 2008; L. Zhang et al., 2010). L. Zhang et al. 125
- (2010) found in particular that the GEOS-Chem simulation for 2006 is unbiased in the 126 middle troposphere at northern mid-latitudes compared with ozonesondes and satellite
- 127 measurements.
- 128
- 129 We use a nested version of GEOS-Chem (Y.X. Wang et al., 2004; Chen et al., 2009; Y.X.
- 130 Wang et al., 2011) with the native  $1/2^{\circ} \times 2/3^{\circ}$  horizontal resolution over North America
- 131 and adjacent oceans (140°-40°W, 10°-70°N), and  $2^{\circ} \times 2.5^{\circ}$  horizontal resolution for the
- 132 rest of the world. We first conduct the global GEOS-Chem simulation at  $2^{\circ} \times 2.5^{\circ}$
- 133 resolution, and then use the output archived at 3-hour temporal resolution as dynamic
- 134 boundary conditions for the nested model.
- 135
- 136 Global anthropogenic emissions are from the Emission Database for Global Atmospheric 137 Research (EDGAR) inventory (Olivier and Berdowski, 2001) for 2000, superseded by
- 138 regional emission inventories from the EPA 2005 National Emission Inventory (NEI-05)
- for the US, Q. Zhang et al. (2009) for Asia in 2006, the European Monitoring and 139
- 140 Evaluation Program (EMEP) for Europe (Vestreng and Klein, 2002), the Canada Criteria
- 141 Air Contaminants (CAC) emission inventory for Canada
- 142 (http://www.ec.gc.ca/pdb/cac/cac home e.cfm), and the Big Bend Regional Aerosol and
- 143 Visibility Observational (BRAVO) emission inventory for Mexico (Kuhns et al., 2005).
- 144 The EMEP, CAC, and BRAVO emissions are scaled on the basis of energy statistics to
- 145 2006 as described by van Donkelaar et al. (2008). We doubled the Japanese and Korean
- 146  $NO_x$  anthropogenic emissions from Q. Zhang et al. (2009) as constrained by OMI
- 147 tropospheric NO<sub>2</sub> column measurements (L. Zhang et al. 2008). Anthropogenic NO<sub>x</sub>
- 148 emissions from fertilizer application are from Yienger and Levy (1995).
- 149
- 150 Natural sources of ozone include open fires, lightning, the biosphere, and transport from 151 the stratosphere. We use monthly biomass burning emissions from the Global Fire 152 Emission Database version 2 (GFED-v2) (van der Werf et al., 2006). Lightning  $NO_x$
- 153 emissions are linked to deep convection following the parameterization of Price and Rind
- 154 (1992) with vertical profiles from Pickering et al. (1998). The global spatial distribution
- 155 of lightning flashes is rescaled to match the 10-year climatology of OTD/LIS satellite
- 156 observations (Sauvage et al., 2007) with higher NO<sub>x</sub> yield per flash at northern mid-
- 157 latitudes than in the tropics (Hudman et al., 2007). The global lightning source is imposed
- to be 6 Tg N  $a^{-1}$  (Martin et al., 2007). Soil NO<sub>x</sub> emissions are computed using a modified 158 159
- version of the Yienger and Levy (1995) algorithm with canopy reduction factors as
- 160 described in Y. Wang et al. (1998). Stratospheric ozone is simulated with a linearized

161 ozone (Linoz) parameterization (McLinden et al., 2000) that provides a mechanistic

- 162 representation of stratospheric influences on tropospheric ozone.
- 163

164 We conducted three-year GEOS-Chem simulations for 2006-2008. The standard

simulation includes all sources and is used for evaluation with observations. We also

166 conducted simulations with: (1) zero North American anthropogenic emissions (North
 167 American background or PRB) for 2006-2008, (2) zero US anthropogenic emissions (US
 168 herbergend) for 2006 and (2) zero enthropogenic emissions (US

background) for 2006, and (3) zero anthropogenic emissions worldwide and methane set
 to its 700 ppby pre-industrial value (natural background) for 2006. All ozone

170 concentrations presented in this paper are daily 8-h average maxima (daily 8-h max), the

171 metric used for the US NAAQS. We find that the US background is on average 1-3 ppbv

172 higher than the North American background, reflecting anthropogenic sources in Canada

and Mexico, with little variability except in border regions. Our results for the US

background are similar to those reported in the focused GEOS-Chem analysis of H.

175 Wang et al. (2009) and hence we do not discuss them further.

176

## 177 **3. Evaluation with observations in the intermountain West**

178

179 We evaluated our GEOS-Chem simulation with the nationwide ensemble of surface 180 ozone observations from the Clean Air Status and Trends Network (CASTNet; 181 http://www.epa.gov/castnet), which monitors air quality in rural areas. The CASTNet 182 sites are shown in Figure 1. Comparison results are generally consistent with the previous 183 GEOS-Chem evaluations presented by Fiore et al. (2003) and H. Wang et al. (2009); time 184 series for individual sites and summary statistics for each region are given in the 185 Supplementary Materials. We focus here on the 12 sites in the intermountain west US, identified in Figure 1 and listed in Table 1, for the year 2006. Interannual variability for 186 187 seasonal mean concentrations at individual sites is weak during 2006-2008 in both model 188 and observations, generally less than 2 ppby. However, interannual variation in the 189 number of exceedance days for thresholds of 65-75 ppbv is larger as discussed in Jaffe 190 (2011).

191

192 Figure 2 shows the time series of observed vs. simulated daily 8-h max ozone

193 concentrations in spring-summer 2006 at four representative sites in the intermountain

194 West, and Figure 3 shows scatterplots for the ensemble of sites. Spring-summer is when

195 concentrations are highest. Also shown in the figures are the North American

background (PRB) and natural background values. Mean values and correlation

197 coefficients for the simulated vs. observed ozone time series at all 12 sites are

summarized in Table 1. We find that seasonal mean ozone concentrations in the model

are generally within  $\pm 2$  ppbv of the observations in Table 1. The correlation coefficients between model and observations are only 0.2-0.5 for the individual sites, which may

between model and observations are only 0.2-0.5 for the individual sites, which may reflect the small dynamic range of variability in the observations. The correlation

202 coefficient is 0.6 in spring and 0.3 in summer for the ensemble of sites in Figure 3.

203

The North American background (PRB) averages 39-44 ppbv (spring) and 35-45 ppbv

205 (summer) for the ensemble of sites in the intermountain West and drives most of the day-

206 to-day variability. The North American anthropogenic enhancement (difference between

207 standard and PRB simulations) averages only 10-22 ppby depending on the site. As 208 shown in Figure 3, PRB increases with increasing ozone concentration in the 209 intermountain West, whereas for surface sites in the East there is little correlation of PRB 210 with ozone (see Supplementary Materials). The natural background is on average 25-28 211 ppbv (spring) and 25-33 ppbv (summer) and is strongly correlated with the PRB. The 212 model difference between the PRB and natural ozone reflects intercontinental pollution 213 influences plus anthropogenic methane. It averages 13-16 ppbv in spring and 11-13 ppbv 214 in summer. Intercontinental pollution influence on ozone is larger in spring because of 215 stronger winds and slower chemical loss (Jacob et al., 1999). Annual 4<sup>th</sup>-highest ozone 216 values at the intermountain West sites are 51-59 ppbv for PRB and 34-45 ppbv for the 217 natural background.

218

219 It is of particular interest to evaluate the ability of the model to reproduce the frequencies 220 of exceedance of proposed air quality standards. Figure 4 shows the simulated vs. 221 observed number of days at individual sites when daily 8-h max ozone concentrations 222 exceed thresholds of 60, 65, and 70 ppbv in spring and summer 2006. The model captures 223 most of the ozone exceedances except for the 65 and 70 ppbv thresholds in spring and for 224 the Mesa Verde site in summer. There is observational evidence that stratospheric 225 intrusions cause high-ozone events in the intermountain West in spring (Langford et al., 226 2009) and these may not be properly represented in the model. The Mesa Verde model 227 overestimate may reflect excessive summer lightning NO<sub>x</sub> emissions over Mexico and 228 the US Southwest.

229

When the ensemble of sites is considered as in Figure 3, we find that the model can
provide an unbiased ozone simulation up to about 75 ppbv but fails to reproduce
exceptional events of higher concentrations. Such events are illustrated in Figure 2 for
Pinedale (80 ppbv) and Rocky Mountain NP (91 ppbv). This may reflect a general
difficulty in Eulerian models to preserve the structure of plumes of dimensions
comparable to the grid resolution (Rastigeyev et al., 2010).

236

# 4. Distribution of background ozone and contribution to pollution episodes.

239 Figure 5 shows the simulated and observed frequency distributions of ozone for the 240 ensemble of CASTNet sites in the US in March-August 2006, separately for low-altitude 241 (< 1.5 km) and high-altitude sites. Also shown are the model frequency distributions for 242 North American (PRB) and natural ozone backgrounds. The model is unbiased in its 243 simulation of the overall distribution. The PRB averages  $27 \pm 8$  ppbv at the low-altitude 244 sites and  $40 \pm 7$  ppbv at the high-altitude sites. The natural background averages  $18 \pm 6$ 245 ppbv at the low-altitude sites and  $27 \pm 6$  ppbv at the high-altitude sites. The difference 246 between PRB and natural background reflects intercontinental pollution influence 247 including anthropogenic methane; it averages 9 ppbv at the low-altitude sites and 13 ppbv at the high-altitude sites. The mean 2006 value of the annual 4<sup>th</sup>-highest daily 8-h max 248 249 ozone is 54 ppbv for PRB and 39 ppbv for the natural background at the ensemble of 250 high-altitude sites, compared with 42 ppbv for PRB and 29 ppbv for the natural 251 background at the low-altitude sites.

253 Our general PRB statistics for the US can be compared to the previous GEOS-Chem 254 studies of Fiore et al. (2003) and H. Wang et al. (2009). H. Wang et al. (2009) found a 255 mean PRB of  $26 \pm 8$  ppbv for summer 2001, whereas we find  $30 \pm 10$  ppbv for summers 256 2006-2008. Fiore et al. (2003) reported a typical PRB range of 15-35 ppbv for March-257 October 2001 using a slightly different metric (mean afternoon concentrations). Our 258 results are overall about 4 ppby higher than these previous estimates. A contributing factor is our higher lightning NO<sub>x</sub> source, 6 Tg  $a^{-1}$  as compared to 4.5 Tg N  $a^{-1}$  in H. 259 260 Wang et al. (2009). Another factor is the 2001-2006 increase in Asian anthropogenic  $NO_x$ 261 emissions, which we previously estimated to have increased PRB ozone by up to 3 ppbv 262 in the West in spring (L. Zhang et al., 2008). We also find some dependence on the 263 model resolution, as our outer nest with  $2^{\circ} \times 2.5^{\circ}$  horizontal resolution yields mean PRB 264 values that are 1-2 ppby lower than the nested simulation. Our results are consistent with 265 those of Parrington et al. (2009), who found a 5 ppby increase in background ozone in the 266 western US compared to Fiore et al. (2002) after assimilation of TES satellite ozone data 267 into the GEOS-Chem model.

268

269 Figure 6 shows the spatial distribution of the seasonal mean PRB ozone concentrations 270 for spring and summer 2006. The PRB is highest in the intermountain West because of 271 the combination of high elevation, deep boundary layer mixing, large-scale subsidence, 272 slow ozone deposition to the arid terrain, and slow ozone chemical loss due to dry 273 conditions (Fiore et al., 2002). The PRB ozone generally decreases from spring to 274 summer, reflecting faster chemical ozone loss; this seasonal decrease is particularly 275 pronounced in the Northeast and on the West Coast. An increase in PRB from spring to 276 summer is found in the Southwest due to summer lightning. The maximum PRB value 277 over New Mexico in summer reflects intense lightning and deep boundary layer mixing. 278

279 Also shown in Figure 6 is the mean PRB ozone on the days when simulated daily 8-h 280 max ozone exceeds 60 ppby. We find that the mean PRB on these high-ozone days is 281 higher than the seasonal mean almost everywhere. The difference is particularly 282 pronounced in the West, where the PRB is on average 7 ppbv higher than the seasonal 283 mean for both spring and summer. It is also large in the Great Lakes region in spring 284 where high ozone values are associated with model lightning. Fiore et al. (2002) 285 previously found PRB values to be maximum for ozone concentrations in the 50-70 ppby 286 range, and this is consistent with our result. If the NAAQS is lowered in the 60-70 ppbv 287 range, areas of the intermountain West will have little or no ability to reach compliance 288 through North American regulatory controls.

289

Finally, we show in Figure 7 the simulated annual 4<sup>th</sup>-highest North American background (PRB) ozone in surface air averaged over 2006-2008, representing the lowest air quality standard that can be achieved by North American emission controls. Values are typically 35-45 ppbv in the East and on the West Coast but 50-60 ppbv in the intermountain West, with a maximum of 64 ppbv over New Mexico and a secondary maximum of 59 ppbv over Idaho due to large wildfires in 2007. A recent study with the

296 CMAQ regional model found much larger contributions from wildfires on surface ozone

in the western US (Mueller and Mallard, 2011). Aircraft observations of California fire

plumes indicate however no significant ozone enhancements unless mixed with urbanpollution (Singh et al., 2010).

300

## 301 5. Conclusions

302

303 We have used the GEOS-Chem global 3-D model of atmospheric composition with  $1/2^{\circ}$ 

 $2/3^{\circ}$  nested horizontal resolution over North America to provide updated estimates of the PRB ozone for the US in 2006-2008. Our work is intended to assist the US EPA in its current risk and exposure assessments as part of the NAAQS-setting process.

307

308 We evaluated the GEOS-Chem simulation with the ensemble of ozone observations from 309 CASTNet sites across the US. Comparisons show in general similar results to previous 310 GEOS-Chem PRB studies (Fiore et al., 2003; H. Wang et al., 2009) and are documented 311 in the Supplementary Materials. We focused our attention on the intermountain West, 312 where the PRB is particularly high and may interfere with the achievability of ozone air 313 quality standards. We showed that the model gives an unbiased representation of ozone in 314 that region and that the PRB drives most of the ozone variability. The model captures the 315 frequency of high-ozone events up to about 70 ppbv but fails to reproduce events of 316 exceptionally high ozone that may be due to stratospheric or wildfire influences. We 317 expect following Rastigeyev et al. (2009) that Eulerian models in general would have 318 difficulty in capturing exceptional events.

319

320 We obtained mean PRB values for the US in spring-summer of  $27 \pm 8$  ppby at low-321 altitude sites (< 1.5 km) and  $40 \pm 7$  ppbv at high-altitude sites. These values are 9-13 322 ppby higher than the natural background due to intercontinental pollution including 323 anthropogenic methane. Our PRB estimates are on average 4 ppby higher than in 324 previous GEOS-Chem studies (Fiore et al., 2003; H. Wang et al., 2009) and we attribute 325 this to a combination of increasing Asian emissions, higher model lightning, and higher 326 model resolution. We find that the PRB generally decreases from spring to summer 327 except in regions strongly affected by summer lightning. We also find that the PRB is 328 higher than average when ozone exceeds 60 ppby, particularly in the intermountain West. 329 The annual 4<sup>th</sup>-highest PRB value in the model (representing the minimum standard 330 achievable through suppression of North American anthropogenic emissions) is typically 331 in the 35-45 ppby range over the East and the West Coast but 50-60 ppby in the 332 intermountain West. Such high PRB values in the intermountain West compared to the 333 proposed revisions of the ozone NAAQS (60-70 ppbv) suggest that special consideration 334 of that region may be needed in the NAAQS-setting process.

335

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339

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549 Figure captions

**Figure 1.** CASTNet ozone monitoring sites in the continental United States for 2006.

- Sites in the intermountain West (Table 1) are indicated in red. Pluses denote sites above1.5 km altitude.
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Figure 2. March-August 2006 time series of daily 8-h max ozone concentrations at four
representative sites in the US intermountain West. Model results (red line) are compared
with observations (black line). Also shown is the North American background or PRB
(blue line) and the natural background (green line). The mean concentrations for the time
period and annual 4<sup>th</sup>-highest ozone values (in parentheses) in ppbv are shown inset.

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Figure 3. Simulated vs. observed daily 8-h max ozone concentrations for spring (March-May) and summer (June-August) 2006 at the 12 intermountain West CASTNet sites of
 Table 1. Also shown is the 1:1 line. The box-and-whisker plots (minimum, 25<sup>th</sup>, 50<sup>th</sup>, 75<sup>th</sup>
 percentile, and maximum) give statistics of the North American background (PRB) and
 natural background for 10-ppbv bins of observed ozone concentrations.

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Figure 4. Simulated (GEOS-Chem) vs. observed number of days with daily 8-h max
ozone concentrations exceeding thresholds of 60, 65, and 70 ppbv in spring and summer
2006 at the 12 CASTNet sites in the intermountain West (Table 1). Symbols identify the
individual sites.

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Figure 5. Frequency distributions of daily 8-h max ozone concentrations in MarchAugust 2006 for the ensemble of low-altitude (<1.5 km) and high-altitude CASTNet sites</li>
in the US (Figure 1). Model results (red) are compared to observations (black). Also
shown are frequency distributions for the North American background (solid blue) and
natural background (dashed green).

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Figure 6. North American background (PRB) ozone concentration in surface air for
spring and summer 2006. The top panels show seasonal means while the bottom panels
show the means for days with ozone > 60 ppbv. Gray areas in the bottom panels had no
days with ozone > 60 ppbv.

- 581
- 582 **Figure 7.** Annual 4<sup>th</sup>-highest value of North American background ozone (PRB)
- calculated in GEOS-Chem as daily 8-h max and averaged for 2006-2008.
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Sites <sup>b</sup>	r	Spring		Summer		Annual 4 <sup>th</sup> highest	
		Observed	<b>GEOS-Chem (PRB)</b> <sup>c</sup>	Observed	GEOS-Chem (PRB)	Observed	GEOS-Chem (PRB)
<b>Yellowstone N.P., WY</b> (44.6N, 110.4W, 2.4 km)	0.40	56.7	52.7 (40.8)	56.0	47.2 (35.3)	69.6	61.4 (51.3)
<b>Pinedale, WY</b> (42.9N, 109.8W, 2.4 km)	0.48	56.5	54.8 (41.9)	57.3	54.1 (38.6)	68.4	65.7 (53.5)
<b>Centennial, WY</b> (41.4N, 106.2W, 3.2 km)	0.20	59.5	54.9 (42.4)	56.1	56.1 (40.1)	70.4	66.6 (52.9)
<b>Rocky Mountain NP, CO</b> (40.3N, 105.6W, 2.8 km)	0.33	56.5	57.9 (44.6)	59.8	61.7 (39.4)	76.1	77.4 (55.3)
<b>Gothic, CO</b> (38.9N, 107.0W, 2.9 km)	0.21	58.8	56.0 (44.4)	53.9	55.4 (40.0)	70.0	65.3 (55.5)
<b>Mesa Verde N.P., CO</b> (37.2N, 108.5W, 2.2 km)	0.31	58.5	57.9 (44.4)	61.2	68.3 (45.6)	74.4	79.6 (58.1)
<b>Great Basin N.P., NV</b> (39.0N, 114.2W, 2.1 km)	0.52	54.5	52.8 (41.2)	58.9	59.2 (40.6)	72.2	72.6 (52.6)
<b>Canyonlands N.P., UT</b> (38.5N, 109.8W, 1.8 km)	0.36	56.6	56.3 (43.4)	59.7	60.3 (42.1)	70.6	70.9 (56.1)
<b>Grand Canyon N.P., AZ</b> (36.1N, 112.2W, 2.1 km)	0.38	58.8	56.6 (43.8)	58.8	60.6 (42.7)	70.8	69.9 (56.4)
<b>Petrified Forest, AZ</b> (34.8N, 109.9W, 1.7 km)	0.57	56.7	55.4 (42.5)	61.5	61.7 (43.3)	71.5	75.2 (57.9)
<b>Chiricahua NM, AZ</b> (32.0N, 109.4W, 1.6 km)	0.41	54.7	53.8 (43.1)	56.5	61.3 (45.1)	74.0	72.3 (58.9)
<b>Big Bend NP, TX</b> (29.3N, 103.2W, 1.1 km)	0.49	52.4	51.3 (39.3)	48.6	54.6 (40.8)	65.3	65.0 (52.3)

1 **Table 1.** Ozone concentrations at CASTNet monitoring sites in the US intermountain West <sup>a</sup>

2 <sup>a</sup> Seasonal mean and annual 4<sup>th</sup>-highest daily 8-h max ozone values in ppbv for 2006 and correlation coefficients (*r*) between model and

3 observations for the daily data. Spring is March-April and summer is June-August.

4 <sup>b</sup>-NP = National Park, NM = National Monument, WY = Wyoming, CO = Colorado, NV = Nevada, UT = Utah, AZ = Arizona, TX = Texas.

<sup>c</sup> GEOS-Chem values in parentheses are the policy-relevant background ozone (PRB) as determined by a simulation with zero North

6 American anthropogenic emissions.

1 Figures





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Figure 1. CASTNet ozone monitoring sites in the continental United States for 2006. Sites in the intermountain West (Table 1) are indicated in red. Pluses denote sites above 1.5 km altitude.

Rocky Mtn NP, CO (40N, 105W, 2743m)







13 Figure 2. March-August 2006 time series of daily 8-h max ozone concentrations at four 14 representative sites in the US intermountain West. Model results (red line) are compared 15 with observations (black line). Also shown is the North American background or PRB (blue line) and the natural background (green line). The mean concentrations for the time 16 period and annual 4<sup>th</sup>-highest ozone values (in parentheses) in ppbv are shown inset. 17





Figure 3. Simulated vs. observed daily 8-h max ozone concentrations for spring (March-May) and summer (June-August) 2006 at the 12 intermountain West CASTNet sites of
 Table 1. Also shown is the 1:1 line. The box-and-whisker plots (minimum, 25<sup>th</sup>, 50<sup>th</sup>, 75<sup>th</sup>
 percentile, and maximum) give statistics of the North American background (PRB) and
 natural background for 10-ppbv bins of observed ozone concentrations.



Figure 4. Simulated (GEOS-Chem) vs. observed number of days with daily 8-h max
ozone concentrations exceeding thresholds of 60, 65, and 70 ppbv in spring and summer
2006 at the 12 CASTNet sites in the intermountain West (Table 1). Symbols identify the
individual sites.





Figure 5. Frequency distributions of daily 8-h max ozone concentrations in MarchAugust 2006 for the ensemble of low-altitude (<1.5 km) and high-altitude CASTNet sites</li>
in the US (Figure 1). Model results (red) are compared to observations (black). Also
shown are frequency distributions for the North American background (solid blue) and
natural background (dashed green).



- days with ozone > 60 ppbv.



Annual 4<sup>th</sup> highest PRB ozone for 2006-2008

### Supplemental Material





Figure S1. March-August 2006 time series of daily 8-h max ozone concentrations at the
additional intermountain West sites of Table 1. Model results (red line) are compared
with observations (black line). Also shown is the North American background or PRB
(blue line) and the natural background (green line). The mean concentrations for the time
period and annual 4<sup>th</sup>-highest ozone values (in parentheses) in ppbv are shown inset.



Figure S2. The same as Figure S1, but for representative CASTNet sites in the Northeast
 US (top), Great Lakes (middle), and the Southeast US (bottom).





Figure S3. Simulated vs. observed daily 8-h max ozone concentrations for spring
(March-May) and summer (June-August) 2006 for the ensembles of CASTNet sites in the
Northeast US, Great Lakes, and the Southeast US. Also shown is the 1:1 line and North
American background (PRB) and natural background model statistics for 10-ppbv bins of
observed ozone concentrations: the minimum, 25<sup>th</sup>, 50<sup>th</sup>, 75<sup>th</sup> percentile, and maximum.