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Impact of Wildfires on Ozone Exceptional Events in the Western U.S.

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- Supporting Information

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ABSTRACT: Wildfires generate substantial emissions of nitrogen oxides (NO_x) and volatile organic compounds (VOCs). As such, wildfires contribute to elevated ozone (O_3) in the atmosphere. However, there is a large amount of variability in the emissions of O_3 precursors and the amount of O_3 produced between fires. There is also significant interannual variability as seen in median O_3 , organic carbon and satellite derived carbon monoxide mixing ratios in the western U.S. To better understand O_3 produced from wildfires, we developed a statistical model that estimates the maximum daily 8 h average (MDA8) O_3 as a function of several meteorological and temporal variables for three urban areas in the western U.S.: Salt Lake City, UT; Boise, ID; and Reno, NV. The model is developed using data from June-September 2000–2012. For these three locations, the statistical model can explain 60, 52, and 27% of the variability in daily MDA8. The Statistical Model Residual (SMR) can



give information on additional sources of O₃ that are not explained by the usual meteorological pattern. Several possible O₃ sources can explain high SMR values on any given day. We examine several cases with high SMR that are due to wildfire influence. The first case considered is for Reno in June 2008 when the MDA8 reached 82 ppbv. The wildfire influence for this episode is supported by PM concentrations, the known location of wildfires at the time and simulations with the Weather and Research Forecasting Model with Chemistry (WRF-Chem) which indicates transport to Reno from large fires burning in California. The contribution to the MDA8 in Reno from the California wildfires is estimated to be 26 ppby, based on the SMR, and 60 ppby, based on WRF-Chem. The WRF-Chem model also indicates an important role for peroxyacetyl nitrate (PAN) in producing O₃ during transport from the California wildfires. We hypothesize that enhancements in PAN due to wildfire emissions may lead to regional enhancements in O₃ during high fire years. The second case is for the Salt Lake City (SLC) region for August 2012. During this period the MDA8 reached 83 ppbv and the SMR suggests a wildfire contribution of 19 ppbv to the MDA8. The wildfire influence is supported by PM_{2.5} data, the known location of wildfires at the time, HYSPLIT dispersion modeling that indicates transport from fires in Idaho, and results from the CMAQ model that confirm the fire impacts. Concentrations of PM_{2.5} and O₃ are enhanced during this period, but overall there is a poor relationship between them, which is consistent with the complexities in the secondary production of O₃. A third case looks at high MDA8 in Boise ID during July 2012 and reaches similar conclusions. These results support the use of statistical modeling as a tool to quantify the influence from wildfires on urban O_3 concentrations.

I. INTRODUCTION

⁴⁰ Wildfires generate substantial emissions of particulate matter $_{11}$ (PM) and ozone (O₃) precursors. ^{1,2} They are also a major driver for interannual variations in summer air quality in the western U.S. for O₃, ³ PM⁴ and black carbon aerosol. ⁵ However, degree of O₃ production from wildfires is highly variable. In a recent review on O₃ production, the majority of published studies dientified a positive relationship between carbon monoxide (CO) and O₃ in wildfire plumes, which is a good indicator of O₃ production. ⁶ For boreal and temperate fires, the Δ O₃/ Δ CO

ratios were on average 0.018 ± 0.051 , 0.15 ± 0.03 and $0.22 \pm_{49}$ 0.23 ppbv ppbv⁻¹ (± 1 s.d.) for plumes aged 1–2 days, 2–5 $_{50}$ days and ≥ 5 days, respectively, showing that O_3 production $_{51}$ generally increased with age of the plume, but with large plume- $_{52}$ to-plume variability.

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 O_3 production is complicated by a number of factors including highly variable emissions, O_3 aerosol effects; O_4 so for emissions of oxygenated O_4 VOCs^{1,10,11} that can result in rapid conversion of NOx to PAN. If a wildfire plume mixes with urban emissions, more rapid O_3 production than either the fire or urban emissions would generate by themselves is likely. O_4 The complexity of emissions, meteorology, radiation and aerosol effects make it very difficult to accurately model O_3 photochemistry using standard Eulerian chemical models.

For O₃, the U.S. EPA uses the 1 h daily maximum and the 65 maximum daily 8 h average O₃ (MDA8) for its regulatory 66 standards. Across most of the western U.S., background O₃ is 67 already elevated due to the high elevations and exposure to the 68 free troposphere. 14-17 During high fire years the distribution of 69 MDA8 values across the western U.S. shifts by 5-7 ppbv, 70 making compliance with the O₃ standard much more 71 challenging. 14 For this reason, the EPA has developed a policy 72 on "exceptional events," which can be defined as "Unusual or 73 natural events that affect air quality but are not reasonably 74 controlled..." (see http://www.epa.gov/ttn/analysis/exevents. 75 htm). Exceptional events can include natural dust storms, 76 transport from sources outside of North America, transport of 77 air from the upper troposphere or lower stratosphere or 78 pollution impacts due to wildfires. To exclude data from 79 consideration, a region must submit a request to EPA that 80 demonstrates, quantitatively, that the air quality would have 81 met the appropriate standard but for the exceptional event.

One approach to identify exceptional events is to use a regression model that calculates O₃ mixing ratios as a function of various meteorological variables. In most cases, daily maximum temperature has been found to be the best predictor for peak or MDA8 O₃ at most sites. Other studies have included a wide array of meteorological variables, such as temperature, cloud cover, or humidity to improve the model fit. In one study of 74 regions in the eastern U.S., In oddifferent variables were considered in the model. Daily maximum temperature and daily average relative humidity (RH) were found to be the most important predictors for MDA8 O₃. Using a generalized additive model, the authors were able to predict the MDA8 values with an R² of 0.5–0.7 for most sites.

In this paper we demonstrate a statistical model that can predict the MDA8 O_3 for three metropolitan regions in the western U.S.: Salt Lake City, Utah; Boise, Idaho; and Reno, Poevada. For three cases we examine days that have high residuals from the statistical model and propose that these were due to wildfire influence based on a variety of supporting data (e.g., Community Multiscale Air Quality (CMAQ) and Weather Research and Forecasting Model with Chemistry (WRF-CHEM) Eulerian models, satellite data, air mass trajectories, etc.). The residual from this statistical model can provide quantitative information on the O_3 contribution. This work can help guide future analyses to quantify the influence of wildfires on O_3 in urban areas, in support of exceptional event designations.

II. MATERIALS AND METHODS

110 We used data from a variety of sources for this analysis 111 including the EPA's AQS sites, CASTNET O₃ measurements, 112 NPS O₃ measurements, IMPROVE aerosol measurements, 113 meteorological data from NCDC and satellite observations 114 from instruments onboard the NASA A-train constellation.

Gridded meteorological data were from the NCEP/NCAR 115 Reanalysis data set. For the Salt Lake City (SLC) urban area, 116 we used the daily average MDA8 from all AQS sites in Salt 117 Lake and Davis Counties, Utah (up to four sites). For Boise 118 and Reno, we used the daily average of all MDA8 values in each 119 metropolitan statistical area as defined by EPA (up to three and 120 six sites, respectively). We also used surface meteorological data 121 from the SLC, Boise and Reno airports. We used satellite data 122 for CO mixing ratios derived for 800 mb from the AIRS 123 instrument onboard the Aqua satellite and aerosol optical depth 124 (AOD) from the MODIS instruments on board the Aqua and 125 Terra satellites. When daily data from both instruments were 126 available, we averaged the AOD values. For all analyses, we 127 used data for the primary fire season, June-September 2000- 128 2012, except there is no data for Boise in 2000. Further details 129 and data sources are given in Table S1 in the Supporting 130 Information.

We used two different Eulerian models to help quantify the 132 contribution to the MDA8 from wildfires, since neither model 133 had results for the full time period of our study. For the 2012 134 wildfires, we used the CMAQ model²² to quantify the influence 135 of wildfires on O₃ concentrations. CMAQ is a peer-reviewed, 136 state-of-the-science Eulerian photochemical model that is run 137 daily as part of the experimental BlueSky Gateway air quality 138 modeling system, which quantifies air pollutant concentrations 139 resulting from wildfires and other emissions sources on a 140 national scale.²³ BlueSky Gateway combines meteorological 141 predictions from the Pennsylvania State University/National 142 Center for Atmospheric Research Mesoscale Model (MM5) 143 version 3.7 with air quality predictions from CMAQ version 144 4.5.1 at a coarse (36 km) grid resolution, and aerosol tracers 145 have been implemented in CMAQ to track primary PM_{2.5} 146 generated by fires. Fire emissions in the CMAQ simulations 147 were derived from fire activity data (locations and acres 148 burned) and the SMARTFIRE system. Fuel loadings were 149 derived from the Fuel Characteristic Classification System 150 (FCCS), fuel consumption is estimated by the Consume 3.0 151 model, and fire emissions are calculated using the Fire Emission 152 Production Simulator (FEPS). The emissions for each fire 153 event vary by day and hour as the fire progresses. For the 154 summer of 2012, CMAQ was run with and without fire 155 emissions to evaluate fire impacts on O₃ concentrations across 156

For 2008, we performed simulations with the regional WRF- 158 Chem version 3.2²⁴ to quantify the influence from wildfires. 159 The model covers the contiguous U.S. at a horizontal 160 resolution of 24 × 24 km and is run for the time period 161 from 10 June to 10 July 2008. The anthropogenic emissions are 162 obtained from the U.S. EPA 2005 National Emissions 163 Inventory (NEI-2005). Biomass burning emissions are obtained 164 from the Fire INventory from NCAR (FINN V1)²⁵ and are 165 distributed in the model vertically following the online plume- 166 rise module.²⁶ The model is configured for the MOZART gas 167 phase chemical scheme linked to the GOCART aerosol 168 model.²⁷ A more detailed description of the model config- 169 uration can be found in ref 28.

In addition to standard chemical tracers we include in these 171 model runs a synthetic tracer that keeps track of O_3 that is due 172 to NO_x emissions from fires. The O_3 tracer method or "XNO_x" 173 method is described in detail in ref 29 and has been used in 174 global models for identifying source contributions, such as for 175 quantifying the O_3 budget $^{30-34}$ and here for the first time it has 176 been applied in a regional model. The method tags emissions of 177

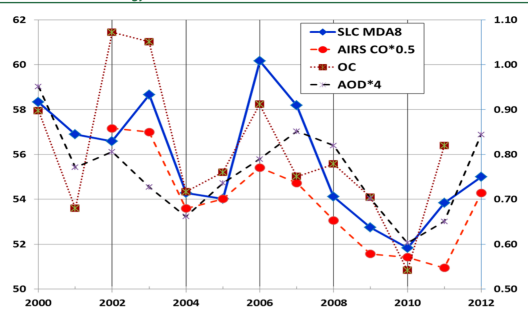


Figure 1. Time series of summer (June–September) median MDA8 (ppbv, left axis) from Salt Lake City urban area. Also shown are summer median organic carbon (OC, μ g/m³, right axis) from three background IMPROVE sites (CRM, PND, GRB) and two satellite observations: CO and AOD. Units are ppbv (left axis) for SLC MDA8, ppbv*0.5 (left axis) for the AIRS CO data, μ g m⁻³ (right axis) for OC. For AOD, the value is multiplied by 4 and shown on the right axis. The correlation coefficient between the annual median SLC MDA8 and OC, AIRS CO and AOD are 0.75, 0.86, and 0.58, respectively. For satellite data (CO and AOD), these are averaged over 38–43° N -110-115° W.

Table 1. Regression Models for SLC, Boise and Reno MDA8 (ppbv)^a

	unstandardized coefficients		standardized coefficients		
SLC model	В	std. error	beta	t	sig.
constant	814	95.8		8.50	0.00
daily max temp	1.29	0.035	0.634	36.7	0.00
daily avg. wind spd.	-0.197	0.014	-0.237	-13.7	0.00
Yr	-0.388	0.048	-0.132	-8.14	0.00
DOY^2	-1.95×10^{-4}	1.24×10^{-5}	-0.256	-15.6	0.00
700 mb zonal wind	-0.615	0.076	-0.140	-8.06	0.00
unstandardized coefficients		coefficients	standardized coefficients		
Boise model	В	std. error	beta	t	sig.
constant	781	111		7.03	0.00
daily max temp	1.02	0.036	0.562	28.1	0.00
daily avg. wind spd.	-0.087	0.019	-0.087	-4.59	0.00
yr	-0.372	0.055	-0.123	-6.74	0.00
DOY^2	-1.85×10^{-4}	1.34×10^{-5}	-0.260	-13.8	0.00
700 mb zonal wind	-0.521	0.068	-0.151	-7.66	0.00
	unstandardized coefficients		standardized coefficients		
reno model	В	std. error	beta	-	sig.
constant	32.6	1.49		21.9	0.00
daily max temp	0.813	0.039	0.456	20.7	0.00
DOY^2	-1.11×10^{-4}	1.27×10^{-5}	-0.193	-8.76	0.00
700 mb zonal wind	-0.189	0.059	0.071	-3.20	0.00

[&]quot;Overall R^2 for the SLC, Boise and Reno models are 0.60, 0.52, and 0.27, respectively. Units for MDA8, $T_{\rm max}$ and wind speeds are ppbv, °C and m sec⁻¹, respectively. Inclusion of previous day's MDA8 increases the R^2 values to 0.65, 0.57, and 0.47, respectively. Note that while DOY is initially included as a quadratic term, there is no difference in the final model fit by including it as a squared term.

178 NO and its resulting nitrogen-containing products (e.g., HNO₃, 179 PAN, HNO₄, etc.) and follows them to the production of O₃. 180 In addition to the standard tagging method, we further 181 conducted a simulation where we did not allow O₃ to be 182 produced through PAN decomposition from fires, in order to 183 provide an estimate of the role of PAN on O₃ production in fire 184 plumes.

III. RESULTS

As noted above, there are large interannual variations in the $_{185}$ summer mean MDA8, with a range of 49–58 ppbvfor SLC.To $_{186}$ examine the relationship between seasonal MDA8 and other $_{187}$ parameters that are likely associated with wildfires, we $_{188}$ compared the SLC and regional MDA8 with organic carbon, $_{189}$ AIRS CO and AOD. Figure 1 shows this comparison using $_{190}$ fi

191 summer median values for 2000-2012. In all cases, the median 192 values are significantly correlated ($p \le 0.05$) with SLC median 193 MDA8. Thus it is reasonable to propose that wildfires in the 194 western U.S. may be the primary driver to explain these large 195 interannual variations and may cause a significant shift in 196 median MDA8, up to +9 ppbv in SLC. While this broad 197 seasonal comparison does not help identify wildfire impacts on 198 individual days, it does demonstrate the challenge regions have 199 in meeting the O₃ air quality standard during high fire years. 200 Also apparent in Figure 1 is a downward trend in median 201 MDA8 values in the SLC region. This is likely due to regional 202 emissions controls, as demonstrated by the downward trend in 203 urban NO_x concentrations for the SLC region. NO_x 204 concentrations in the region have decreased approximately 205 5% per year since 2000 (see http://www.epa.gov/airtrends/ 206 nitrogen.html).

Statistical Model Development. We used PASW 207 208 Statistics software (now IBM SPSS Statistics) version 18.0.3. 209 to develop the statistical model. For each location, we examined 210 the multilinear relationship (MLR) between the indicated 211 variable and the regional averaged daily MDA8 value for June-212 September. A large number of variables were considered to 213 identify the best model including surface variables (daily 214 maximum temperature, daily average wind speed) and upper air 215 parameters (see Table S1). We identified the variables that gave 216 the strongest explanatory power (correlation) with the least 217 multicollinearity, as this would confuse the interpretation of the 218 model results.³⁵ For all locations, we found that daily maximum 219 temperature was the strongest predictor for MDA8. Supporting 220 Information Figure S1 shows a scatter plot of MDA8 versus 221 daily maximum temperature for SLC. However, other variables 222 also show a significant relationship with MDA8 and should be 223 included in the model. The MLR model fits an equation of the 224 form

$$MDA8 = a + bX_1 + cX_2... + residual$$

225 Where the MDA8 is the dependent variable, X_1 , X_2 , etc. are the 226 independent predictors, a, b, c, etc. are the coefficients and the 227 residual is the unfit portion of the model. In this analysis we 228 refer to the residual as the statistical model residual (SMR). We 229 examined the best form for each predictor including linear, 230 squared, quadratic, log, etc. For most variables a linear fit gave 231 the best performance, except for day of year, where a squared 232 term yielded an improved fit.

We also examined the performance of generalized linear models (GLM) as a tool to build the statistical models, but did not find a significant improvement in predictive ability. Therefore we used the simpler MLR approach, which makes interpretation of the predictors more straightforward.

Table 1 summarizes the model predictors and Table 2 provides a summary of the daily MDA8 and daily model

Table 2. Statistical Summary of SLC, Boise, and Reno MDA8 and Model Residuals (June-September)

	N	minimum	maximum	mean	std. deviation
SLC MDA8	1585	19.7	101.5	55.8	11.0
SLC Resid.	1582	-24.7	34.3	0.0	7.0
Boise MDA8	1449	17.0	86.0	50.9	10.3
Boise Resid.	1449	-30.8	26.1	0.0	7.2
Reno MDA8	1586	24.8	82.0	52.8	8.4
Reno Resid.	1586	-27.7	25.9	0.0	7.2

residuals for each location. For SLC and Boise, the best fit 240 model explained 60 and 52% of the variance in MDA8 values. 241 For these two regions we found that daily maximum 242 temperature, daily average surface wind speed, day of year², 243 year and the 700 mb zonal wind component gave the best fit 244 with minimal multicollinearity. Inclusion of additional mete- 245 orological variables made virtually no difference to the model 246 fit. Expanding the analysis to include May data generally 247 reduced the performance of the model in all three locations. 248 For Reno, the model explained less of the total variance (27%) 249 and the variables included only daily maximum temperature, 250 day of year² and the 700 mb zonal wind component. The most 251 likely explanation for this is that SLC and Boise, are more 252 isolated cities, whereas Reno is adjacent to larger emissions 253 sources in California. We expect that a parameter that included 254 a better measure of transport from California might explain 255 more of the variance in MDA8 values.

Because the MDA8 values have significant autocorrelation, 257 inclusion of the previous day's MDA8 value will improve the 258 model fit. For SLC and Boise, this improvement was relatively 259 minor (R^2 values increased by 0.05 for each), whereas for Reno 260 the model fit was improved substantially (R^2 increased by 0.20). 261 For most days, this had relatively little impact on the calculated 262 residual, except in a multiday pollution event. For such an 263 event, the residual for the second and succeeding days was 264 reduced if the previous day's MDA8 was included as a model 265 predictor. However, since our goal is to identify exceptional 266 events, we feel that these should be predicted from 267 meteorological parameters alone, not the previous day's 268 MDA8 values. Thus the final statistical model used in this 269 analysis does not include the previous day's MDA8 as a 270 predictor. As a result, the SMRs show significant autocorrela- 271 tion. We examined the influence that this autocorrelation has 272 on the SLC model results as follows. We removed 80% of the 273 data points by selecting data on only every fifth day and then 274 reran the model with the same predictors. We found that this 275 made very little difference in the overall R^2 of the model and all 276 predictor variables remained statistically significant. The 277 residuals from this reduced data model (every fifth day) no 278 longer show significant autocorrelation. With the reduced data 279 set, the Durbin-Watson test confirmed no autocorrelation at a 280 significance level of p = 0.05 (test statistic = 1.93), 281 demonstrating that autocorrelation has minimal influence in 282 the reduced data model. So, while autocorrelation is present in 283 the residuals for the full data model, we found it has little 284 influence on the predictors or the form of the model.

It is important to check the distribution of the residuals and 286 the relationship with respect to the original independent 287 variables. A histogram of the SMR values for SLC is shown in 288 Supporting Information Figure S2 and a plot of SMR versus 289 daily maximum temperature is shown in Supporting 290 Information Figure S3. Mean and standard deviation (SD) 291 for the SMRs for each location is shown in Table 2. At all 292 locations, the SMR has a mean of 0 and a SD that is smaller 293 than the SD of the MDA8. The residuals are normally 294 distributed and show no pattern with respect to temperature. 295 Similar results are found when the residuals are plotted against 296 the other independent variables in the model for all locations. 297

Interpretation of the Statistical Model Residual (SMR). 298 The SMR can yield information about days that have higher 299 MDA8 than predicted from the input meteorological 300 conditions. These days are then candidates to consider as 301 exceptional events, but further evidence is needed to identify 302

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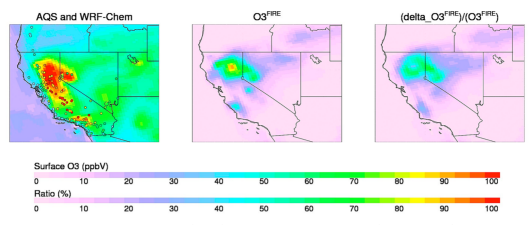


Figure 2. Surface O_3 from AQS sites for 24 June 2008 (circles) and WRF-Chem model results (ppbv, left panel), O_3 due to fires in WRF-Chem (ppbv, middle panel) and % of O_3 from fires where the NO_x cycled through PAN decomposition (%, right panel). Observations are an average for hours 15–17 local time, model results for 0 UTC (~16 local time).

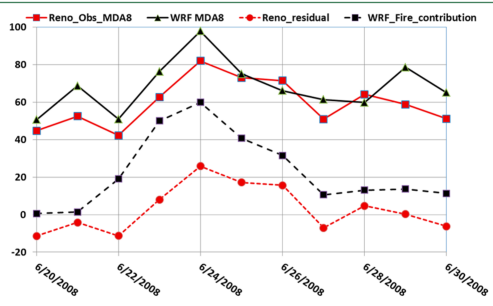


Figure 3. Observed MDA8 and residual from the statistical model for Reno. Calculated MDA8 from the WRF model and the model calculated contribution from fires.

 $_{303}$ the cause of high $_{O_3}$. Possible causes for high SMR might $_{304}$ include

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- Additional precursors from unusual sources within the regio;
- (2) Additional O₃ produced from precursors emitted by wildfires;
- 309 (3) Unusually large contribution of O₃ from the upper troposphere/lower stratosphere (UTLS);
 - (4) Unusually large contribution from transport of O₃ and/ or precursors from distant sources.

Both transport from Asian sources and transport from the 314 UTLS have been previously identified as important sources of 315 O_3 in the western U.S. $^{14,15,36-38}$ Wildfires have also been 316 suggested as important O_3 sources, 6 especially in the western 317 U.S. 3,6,34 To examine the utility of the SMR as a tool to 318 quantify the influence on specific days, we will focus here on 319 three cases with high SMR.

While 2008 was not an exceptional year over the entire western U.S., wildfires in California burned approximately 1.5 million acres in 2008 compared with 0.7 million acres on average for the state between 1997 and 2012 (data from the

National Interagency Fire Center www.nifc.gov). Exceedances 324 of the hourly O₃ standard were reported at a number of sites in 325 California and several of these were considered "exceptional 326 events" by the state.³⁹ Figure 2 shows the modeled O₃ for local 327 f2 afternoon on 24 June 2008. While the fires were located mainly 328 in California, westerly winds carried plumes into Nevada, with 329 O₃ reaching up to ~100 ppbv. A tagged WRF-CHEM model 330 run indicates that between 40 and 60 ppbv of O₃ was 331 contributed by the wildfires across a large section of western 332 Nevada. Figure 3 shows a time series of the measured MDA8 333 f3 and modeled MDA8 in Reno, NV for 20-30 June 2008. The 334 figure also shows the WRF-CHEM fire contribution to the 335 MDA8 and the SMR. Both the measured and modeled MDA8 336 values peak on 24 June, as does the wildfire contribution and 337 the SMR. The MDA8 in Reno on 24 June was 80 ppbv and the 338 SMR suggests that 24 ppbv was due to the wildfire 339 contribution. The calculated fire contribution using WRF- 340 CHEM is 60 ppbv, a much higher value than the SMR; 341 however, we expect this to differ as it shows the tagged 342 contribution under the chemical regime of the fire plume, 343 whereas the SMR is the residual from what is typical for the 344

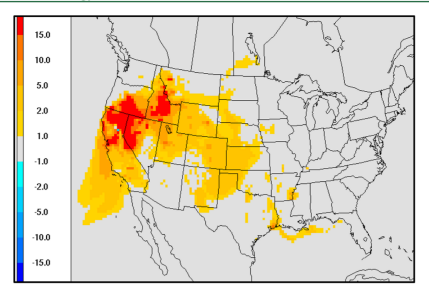


Figure 4. Calculated contribution to MDA8 (ppbv) on 12 August 2012 due to wildfires in the western U.S. This estimate is produced by running CMAQ with and without wildfire emissions. The difference is assumed to be the wildfire contribution. For SLC, the observed MDA8 is 80 ppbv. The modeled MDA8 is 74 ppbv, with a wildfire contribution of 5.5 ppbv. By comparison, the SMR for this day is 19.6 ppbv.

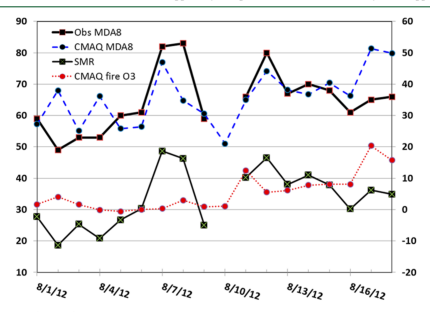


Figure 5. Observed and modeled (CMAQ) MDA8 for SLC for 1–18 August 2012. Also shown is the contribution due to wildfires from the CMAQ model and the SMR. Observations and the SMR are not available for August 10, 2012.

345 meteorological conditions. The consistency in timing between 346 the WRF-CHEM and observed values and the WRF-CHEM 347 wildfire contribution supports the use of the SMR as an 348 indicator of the magnitude of the wildfire contribution. The 349 wildfire influence on 24 June is also supported by Supporting 350 Information Figure S6, which shows the location of fires 351 burning at the time, smoke from the MODIS instrument on the 352 Aqua satellite and information on transport from HYSPLIT 353 trajectories.

Figure 2c also shows that a large fraction of the O_3 that is produced from wildfire precursors is due to NO_x that has been cycled through peroxyacetyl nitrate (PAN). Substantial production of PAN in wildfires has been noted previously, and the WRF-CHEM results demonstrate the importance of PAN in generating O_3 far downwind of the fire region. This may also explain part of the reason O_3 and PM enhancements from some wildfires show little relationship and suggests that

fire influences on O_3 can occur far downwind of the emission $_{362}$ source, driven by PAN transport and decomposition back to $_{363}$ NO...

The 2012 wildfire season was unusually strong across most of 365 the western U.S. In total more than 7 million acres burned in 366 the western U.S. compared to approximately 4 million acres in 367 an average year. Unusually large areas burned in California, 368 Oregon, Idaho, Nevada and Montana in 2012. In late July and 369 early August a large number of fires burned across northern 370 California, southeastern Oregon, northern Nevada and southern Idaho. Figure 4 shows the contribution to the MDA8 due 372 f4 to wildfires for 12 August 2012 as calculated by the CMAQ 373 model. Figure 5 shows the observed and CMAQ-modeled 374 f5 MDA8 for 1–17 August 2012, as well as the CMAQ fire 375 contribution and the SMR. Over the period between 7 and 18 376 August, the average CMAQ wildfire contribution of 8 ppbv is 377 very close to the average SMR of 9 ppbv. However, the CMAQ 378

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379 model tends to underpredict peak MDA8 values in excess of 75 380 ppbv observed on 7, 8, and 12 August. The CMAO model also 381 shows a relatively weak correlation between the observed and 382 modeled MDA8 for this period ($R^2 = 0.21$). On these days, the 383 CMAQ modeled wildfire contribution are also lower than the 384 SMR values. Supporting Information Figure S4 shows the 385 relationship between observed MDA8 and daily average PM_{2.5} 386 for the SLC region during the fire influenced period. For this 387 period, the R² for this relationship is 0.17, whereas in the 388 CMAQ model it is 0.51. Thus we conclude that while the CMAQ model does capture some of the influence from the 390 wildfires during this period, it has difficultly in accurately 391 modeling the observed MDA8. Further evidence for a wildfire 392 influence in SLC on 7 and 8 August is shown in Supporting 393 Information Figures S7 and S8. These figures show the location 394 of fires and smoke (from the NOAA Hazard Mapping System) and demonstrate that the smoke was clearly present in/around SLC on these dates.

For Boise Idaho, results from the statistical model are shown 397 398 in Tables 1 and 2. Supporting Information Figure S5 shows the 399 SMR and CMAQ fire contribution for Boise in July 2012. 400 During this time period, large fires burning in northern 401 California, Oregon and Idaho were influencing air quality across 402 the western states. The MDA8 values in Boise peaked at 74 ppbv on 11 July 2012. During this time period the CMAQ results showed a relatively weak correlation with the observed 405 MDA8 values in Boise (R^2 of 0.12) and the calculated fire 406 contributions were much smaller than the SMR values (see 407 Supporting Information Figure S5). Further evidence for a wildfire influence in Boise on 11 July is shown in Supporting 409 Information Figure S9. This figure shows the location of fires 410 and smoke (from the NOAA Hazard Mapping System) and 411 demonstrates that the smoke was clearly present in/around 412 Boise on this date.

These findings highlight sources of error associated with the BlueSky Gateway CMAQ modeling, which include parametrizations used to solve the atmospheric momentum are equations, spatial grid cell resolution, and uncertainties associated with fire emissions estimates. As in the Reno tase, the timing and magnitude of the observations and the Eulerian model results support the use of the SMR to quantitatively characterize the O_3 production due to wildfires.

IV. DISCUSSION AND SUMMARY

421 Figure 1 demonstrates that wildfires can have a significant 422 influence on MDA8 levels in urban areas of the western U.S., 423 but quantifying the daily impact is a challenge. Development of 424 a statistical model for O₃ is an important and useful exercise 425 that can indicate the types of meteorological conditions that are 426 conducive to O₃ formation in a specific region. Outliers from 427 this model, called the SMR, can then indicate unusual sources 428 of O₃ or unusual conditions that may qualify as exceptional 429 events per the EPA definitions. For cases where corroborating 430 analyses point to the influence of fire emissions on elevated O₃ 431 concentrations, the SMR can provide an estimate of this 432 impact. This may then satisfy EPA's requirement of a 433 quantitative demonstration that O₃ levels would not have 434 exceeded the standard "but for the unusual" event. The 435 statistical modeling technique described in this paper is able to 436 provide this estimate without requiring the resources and 437 expertise needed for complex Eulerian photochemical modeling 438 of wildfire impacts.

We have estimated the magnitude of the wildfire impact on 439 MDA8 for three cases using the SMR and two different 440 Eulerian models (WRF-Chem and CMAQ). None of these 441 calculations can be considered exact, nor can they be 442 considered identical. Each method gives an estimate of the 443 true wildfire contribution for the given case. The SMR value 444 likely underestimates the true impact. This is because the value 445 is calculated as the outlier, and thus ignores any background or 446 average wildfire contribution that is embedded in the seasonal 447 cycle or relationship with temperature. For the CMAO 448 calculation, the wildfire contribution is calculated as the 449 difference in model runs with all wildfires emissions turned 450 on/off. Using WRF-Chem, the contribution is calculated by 451 tagging each emission source and using this to quantify the O3 452 production. The WRF-Chem results also demonstrate an 453 important role for PAN in redistributing primary wildfire 454 emissions and enhancing O3 over a larger region. One 455 consequence of PAN chemistry and complex aerosol effects is 456 that O₃ and aerosol enhancements associated with wildfires 457 show minimal correlation. 458

ASSOCIATED CONTENT

S Supporting Information

This material is available free of charge via the Internet at 461 http://pubs.acs.org. 462

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Notes

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