

Maximum ozone concentrations in the southwestern US and Texas: Implications of growing predominance of background contribution

David D. Parrish¹, Ian C. Faloona^{2,3}, and Richard G. Derwent⁴

¹ David.D.Parrish, LLC, 4650 MacArthur Ln, Boulder, Colorado, USA

² Air Quality Research Center, University of CA, Davis, CA, USA

³ Department of Land, Air, & Water Resources, University of CA, Davis, CA, USA

⁴ rdscientific, Newbury, Berkshire, UK

Correspondence to: David D. Parrish (david.d.parrish.llc@gmail.com)

10 **Abstract.** We utilize a simple, observational-based model to quantitatively estimate the US anthropogenic, background and
wildfire contributions to the temporal and spatial distributions of maximum ozone concentrations throughout the southwestern
US, including Texas and parts of California. The very different temporal variations of the separate contributions provide the
basis for this analysis: over the past four decades the anthropogenic contribution has decreased at an approximately exponential
rate by a factor of ~6.3, while the US background concentration rose significantly through the 1980s and 1990s, reached a
15 maximum in the mid-2000s, and has since slowly decreased. We primarily analyze ozone design values (ODVs), the statistic
upon which the US National Ambient Air Quality Standards (NAAQS) are based. The ODV is an extreme value statistic that
quantifies the relatively rare maximum observed ozone concentrations; thus, ODV time series provide spatially and temporally
resolved records of maximum ozone concentrations throughout the country. Recent contributions of US background ozone to
ODVs (primarily due to transported baseline ozone) are 64 to 70 ppb over most of the southwestern US, and wildfires (also
20 generally considered a background contribution) add further enhancements of 2 to 6 ppb in southwestern US urban areas. US
anthropogenic emissions from urban and industrial sectors now produce only relatively modest enhancements to ODVs (less
than ~6 ppb in 2020) outside of the three largest urban areas considered (Dallas, Houston and Los Angeles), where the 2020
enhancements were in the 17 to 30 ppb range. As a consequence, US background ozone concentrations now dominate over
US anthropogenic contributions in the western US, including the Los Angeles urban basin, where the largest US ozone
25 concentrations are observed. In the southwestern US, this predominance is so pronounced that the US background plus wildfire
contributions to ODVs approach or exceed the US NAAQS for ozone of 70 ppb (implemented in 2015) and 75 ppb
(implemented in 2008); consequently, NAAQS achievement has been precluded in this region. The large background
contribution in this region has led to a pronounced shift in the spatial distribution of maximum US ozone concentrations; once
ubiquitous nearly nationwide, ODVs of 75 ppb or greater have nearly disappeared in the eastern US, but are still frequent in
30 the southwestern US. By 2021, the trend of maximum ODVs in two of the more highly populated eastern urban areas (i.e.,
New York City and Atlanta) had decreased to the point that they were smaller than those in significantly less populated

southwestern US urban areas, and nearly as small as ODVs recorded at isolated rural southwestern US sites. Two implications arise from these findings. First, alternate emission control strategies may provide more effective approaches to ozone air quality improvement; since background ozone makes the dominant contribution to even the highest observed concentrations, an international effort to reduce northern midlatitude baseline ozone concentrations could be pursued, or a standard based on the anthropogenic increment above the regionally-varying US background ozone concentration could be considered to provide a regionally uniform emission reduction challenge. Second, the predominant contribution of US background ozone across the southwestern US presents a profound challenge for air quality modelling, since a manifold of stratospheric and tropospheric processes occurring on small spatial scales, but over hemisphere-wide distances, must be accurately treated in detail to predict present and future background contributions to daily maximum ozone concentrations at local scales.

1 Introduction and Background

Elevated ambient ozone (O_3) concentrations constitute an air quality issue that has affected many urban areas of the world; in Los Angeles they reached extreme levels, with maximum 1-hr average ozone concentrations exceeding 500 ppb in the mid-1960s (Parrish and Stockwell, 2015). In the US, large reductions in anthropogenic emissions of photochemical ozone precursors, following implementation of air quality improvement policies, substantially lowered urban ozone concentrations throughout the country over the past half century. However, several areas still have not attained the National Ambient Air Quality Standard (NAAQS) for ozone (see <https://www.epa.gov/green-book>). The NAAQS, most recently lowered in 2015, requires that the Ozone Design Value (ODV) at each monitoring site in a region not exceed 70 parts-per-billion (ppb, equal to nmole ozone/mole air). Notably the 2008 NAAQS is still in effect; even though it allows a higher ODV (75 ppb), there are different sets of nonattainment areas designated under the two standards. The ODV is an extreme value statistic defined as the 3-year average of the annual fourth-highest daily maximum 8-hour average (MDA8) ozone concentration. The fourth-highest MDA8 represents ~98th percentile of MDA8 values observed in the warm half of the year when those four highest values generally occur; thus, a time series of ODVs observed at a particular monitoring site is a smoothed record of the temporal evolution of the maximum ozone concentrations impacting that location. Ozone monitoring within the US began in the early 1970s, so ODVs have been collected over nearly five decades at more than 2000 sites throughout the Nation. This observational record reflects detailed, spatially and temporally resolved information regarding the variability of ozone sources and sinks that determine the maximum observed ozone concentrations. Our goal in this paper is to analyze this record to quantify the sources of maximum ozone concentrations in the southwestern US, and to investigate the implications of the results. This region has not previously been analyzed by our approach, and is of particular interest because it is impacted by large background ozone concentrations (e.g., Lin et al., 2012a,b; Zhang et al., 2020). Previous work shows that the background contributions to ODVs in some western US regions exceeds 60 ppb (e.g., Langford et al., 2022), and even has approached 70 ppb, making achievement of the 70 ppb NAAQS quite difficult in those regions (Cooper et al., 2015). The five states - Arizona (AZ), Colorado (CO), Nevada (NV), New Mexico (NM), and Utah (UT) – included in this region have four urban areas – Phoenix AZ, Denver CO,

Las Vegas NV and Salt Lake City UT– that are centers of Marginal or Moderate Ozone Nonattainment Areas (see [US EPA Green Book 8-Hour Ozone \(2015\) Area Information](#), last accessed 27 January 2023); additionally Phoenix AZ and Denver CO are classified as Moderate and Severe-15, respectively, Ozone Nonattainment Areas under the 2008 ozone NAAQS (see [US EPA Green Book 8-Hour Ozone \(2008\) Area Information](#), last accessed 27 January 2023).) We also include Texas in this analysis because it represents a transition region between the southwestern US and the very different Gulf of Mexico region; four Texas urban areas – Dallas, Houston, El Paso, and San Antonio – are centers of Marginal to Moderate Ozone Non-attainment areas designated under the 2015 NAAQS ([US EPA Green Book 8-Hour Ozone \(2015\) Area Information](#), last accessed 31 January 2023); additionally Houston and Dallas are classified as Severe-15 Non-attainment areas under the 2008 ozone NAAQS ([US EPA Green Book 8-Hour Ozone \(2008\) Area Information](#), last accessed 31 January 2023). We examine the ODV time series recorded in these four urban areas, plus five other Texas regions containing smaller cities as well as vast rural areas.

Our analysis is a simple, observation-based quantification based on a conceptual model of the ozone sources that determine ODVs throughout the US. Notably, this approach is very different from the computer-based chemical transport models (CTMs) that traditionally have been used for such quantifications; this paper concludes with Section 5.5 that compares and contrasts these two approaches, and discusses how they complement each other. Our analysis has been applied previously in the western US (Parrish et al., 2017; 2022) and the northeastern US (Parrish and Ennis, 2019). It focuses on the two major contributors to ODVs in both urban and rural areas. The first is the *US background ODV* (i.e., the ODV that would be recorded at a site in the absence of any contribution from US anthropogenic emissions); this is the natural and anthropogenic ozone transported from outside the country, plus any ozone produced within the US from naturally emitted precursors; this quantity provides a direct quantification of the minimum ODV that could be achieved by reducing US anthropogenic precursor emissions alone. The second is the *US anthropogenic ODV enhancement* (i.e., the enhancement of an actual ODV above the US background ODV due to contributions from US anthropogenic emissions); this enhancement is due to photochemical production from US anthropogenic emissions of ozone precursors. These definitions are consistent with the more general term “US background ozone” (USB), which is defined by the US Environmental Protection Agency (e.g., Dolwick et al., 2015) to represent the influence of all sources other than US anthropogenic emissions on a particular ozone concentration. Importantly, neither the US background ODV nor the USB can be directly measured because they are theoretical constructs; the measured ozone concentration in air transported to a location without recent local or regional anthropogenic or continental influences is termed baseline ozone, and may be closely related to USB as will be discussed later. We separately quantify the US background ODV and the US anthropogenic ODV enhancement based on their very different long-term temporal changes. This quantification process fits time dependent functions to tabulated ODV time series. As discussed below, these fitted functions have been designed to accurately represent the known temporal changes of the ODV components. The fitting process yields the values of parameters in those functions, and it is these values that quantify the ODV components. The text box below (which is very similar to that included in Parrish et al., 2022) collects the important terms discussed above with their definitions, and the parameters included in the time dependent functions discussed below.

Deleted :

Important quantitative terms and parameters with definitions

- **MDA8** - Maximum daily 8-hour average ozone concentration at a site on a particular day.
- **Ozone Design Value (ODV)** - 3-year running mean of the annual fourth-highest MDA8 ozone concentration at a site.
- **Baseline ozone** - The ozone concentration measured in air transported to a location without recent local or regional anthropogenic or continental influences, typically derived from observations.
- **US background ozone (USB)** - The ozone concentration that would exist at a time and US location from all sources, except that produced from US anthropogenic emissions, typically derived by zeroing out emissions in chemical transport models.
- **US background ODV** - The ODV that would be recorded at a site in the absence of any contribution from US anthropogenic emissions.
- **US anthropogenic ozone ODV enhancement** - The enhancement of an actual ODV above the US background ODV due to contributions from US anthropogenic emissions.
- a - Fit parameter of Equations 1, 3 and 4 that estimates the US background ODV in year 2000.
- A - Fit parameter of Equation 3 that estimates the US anthropogenic ODV in year 2000.
- τ - Fit parameter of equation 3 that estimates the exponential rate of decrease of the US anthropogenic ozone (or ODV) enhancement with time.

100 Prevailing westerly winds carry midlatitude Pacific marine air into the Western US, and the baseline ozone in that air provides the predominant source of the US background ODV in the western US (Zhang et al., 2020; Parrish et al., 2022). Parrish et al. (2020) show that the temporal change in baseline ozone at northern midlatitudes from 1978 to 2018 is accurately quantified by a quadratic polynomial, where t represents time:

$$\text{baseline } O_3 = a + bt + ct^2, \quad (1)$$

105 We use this same functional form to estimate the temporal variation of the US background ODVs. In this work, the time origin is chosen as the year 2000 (i.e., t equals the year - 2000). Thus, the parameter a , with units ppb O_3 , is the intercept of this function in 2000. The value of this parameter varies with location, and quantifies the magnitude of the baseline ozone concentration in that year. The second and third coefficients in Equation 1 quantify the variability of baseline ozone before and after 2000. Parrish et al. (2020) derived numerical values for these parameters - $b = 0.20 \pm 0.06$ ppb yr^{-1} and $c = -0.018 \pm$
110 0.006 ppb yr^{-2} - that were common to eight northern midlatitude baseline ozone time series measured from surface sites, aircraft and sondes over western Europe and western North America covering altitudes from sea level to 9 km. These same coefficient values well represent 28 published trend analyses of baseline representative data sets collected throughout the western US (Parrish et al., 2021). We use these same coefficient values in this work. The positive value of b and the negative value of c indicates that baseline ozone concentrations increased before 2000, reached a maximum after 2000, and then decreased at later
115 times. Equation 2 gives the year of the maximum of the fitted curve,

$$\text{year}_{\text{max}} = -b/2c + 2000, \quad (2)$$

which is ~ 2006 for the above parameter values.

Parrish et al. (2017) show that the US anthropogenic ODV enhancement in US urban areas has decreased rapidly over the past four decades, a decrease accurately captured by an exponential term with an e -folding time of ~ 22 years; this corresponds
120 to a decrease of a factor of more than 6 from 1980 to 2020. This behavior contrasts markedly with the much smaller changes

that occurred in baseline ozone concentrations. Parrish et al. (2017) and Parrish and Ennis (2019) assumed that baseline ozone remained constant, while Parrish et al. (2022) used Equation 1 to quantify the changes that have occurred in the US background ODV near the US West Coast. Equation 3 defines a simple functional form that captures the physical picture of a US background ODV that slowly varies as do baseline ozone concentrations, plus a rapidly decreasing US anthropogenic ODV enhancement;

$$\begin{aligned} \text{ODV} &= a + bt + ct^2 + A \exp(-t/\tau) \\ &= a + 0.20*t - 0.018*t^2 + A*\exp(-t/\tau), \end{aligned} \quad (3)$$

where the values of the b and c parameters derived by Parrish et al. (2020) have been substituted into the second form of the equation. Here A is the year 2000 magnitude of the exponential term designed to quantify the rapidly decreasing US anthropogenic ODV enhancement, τ is the time constant of that exponential decrease, and t equals the year - 2000 as in Equation 1. An exponential function is chosen to quantify the long-term decrease of US anthropogenic ODV enhancements, because it is mathematically as simple as possible (i.e., has the fewest possible unknown parameters), and successfully accounts for a large fraction of the variance in recorded ODV time series throughout the US (see Parrish et al., 2017; 2022; Parrish and Ennis, 2019); this choice is more fully discussed in Sections S3 and S4 of the Supplement. Regression fits of Equation 3 to ODV time series is the primary basis for analysis in this work; values are derived for the a and A parameters, which provide quantitative estimates of the year 2000 US background ODV and year 2000 US anthropogenic ODV enhancement, respectively. The definitions of the three parameters in Equation 3 are included in the above text box

Parrish et al. (2022) applied the above described analysis to the entire western US coastal region; they quantify a small negative latitude gradient (-0.4 ppb/deg) and a strong positive vertical gradient (~10-15 ppb/km) in the US background ODVs at monitoring sites along the US west coast. The latitude gradient is consistent with earlier model and observational analysis (Zhang et al., 2020; Ziemke et al., 2011). The altitude gradient agrees well with that determined by sondes and aircraft profiles along the California coast (Oltmans et al., 2008; Cooper et al., 2011; Yates et al., 2015; Faloona et al., 2020); however, this vertical gradient is weakened by continental mixing as marine air is advected eastward across the complex terrain of the North American Cordillera. The quantification of such fine scale variations (on the order of 1-2 ppb) provides a sense of the analysis precision, and the agreement with other analyses is evidence that the results of the observation-based quantification are physically realistic. Nevertheless, it is important to draw a careful distinction between the definitions of US background ODV and US anthropogenic ODV enhancement presented above, and their observation-based quantifications that we derive. Since those quantifications rely on the observed temporal changes in ODVs, only sources of precursors effectively controlled by regulatory action can contribute to the derived A parameter value, i.e. the estimated US anthropogenic ODV enhancement. The impact of uncontrolled anthropogenic precursor emissions, such as those from agricultural soils, livestock operations, or volatile chemical products, for example, would potentially contribute to the a parameter value, i.e. the estimated US background ODV. Drawing conclusions from the derived quantifications must carefully consider this issue, and bring in additional considerations to avoid confounding influences. Temporal changes in emissions from wildfires are another potentially confounding influence; Section 3.2 discusses our approach to including these changes in our analysis. Furthermore,

Deleted: consumer

the ODVs represent rare events that may occur at any time during the warm season when the combined USB and anthropogenic ozone contributions maximize. This maximum may not occur on the days when the USB contribution is largest (i.e., the days that determine the US background ODV). Thus, for any given extreme ozone episode the sum of US background ODV and the anthropogenic contribution to that episode may not equal the observed ODV; Section S2 of the Supplement discusses in detail the distinction between the US anthropogenic ODV enhancement and the actual anthropogenic contribution to an ODV.

In this work we extend the observation-based analysis of ODV time series to the southwestern US and Texas; we also survey surrounding states, including an analysis of the distributions of MDA ozone concentrations in California air basins. This quantification of the spatial and temporal variability of the US background ODV is important to improve our understanding of the magnitude and temporal changes of the primary ozone sources, to quantify how the growing dominance of USB has changed the regional distribution of occurrence of ozone air quality standard exceedances, and to discuss the implications for future efforts to improve US ozone air quality. In the following, Section 2 describes the data set that is the basis for this analysis, Section 3 discusses details of the analysis approach, Section 4 presents the results, and Section 5 discusses the conclusions and implications.

2 Data sets analyzed

This work examines two data sets: MDA8 ozone concentrations measured during the 1980–2022 period in four coastal California air basins, and ODVs reported from the beginning of US ozone monitoring through 2021 throughout the US. The California MDA8 ozone concentrations were obtained from California’s Air Resource Board data archive (<https://www.arb.ca.gov/adam/index.html>). The ODVs are computed and published annually by US EPA’s Office of Air Quality Planning and Standards. For each year for each ozone monitoring station in the US, an ODV is calculated if the measurements achieve the specified completeness criteria. All values reported in the focus region were downloaded from the EPA data archive (<https://www.epa.gov/aqs>); only the ODVs marked as valid were retained for analysis. Exceptional events, such as wildfires or stratospheric ozone intrusions, are included in the data set, although in principle they can be removed from the MDA8 monitoring record through an EPA concurrence process as uncontrollable “Exceptional Events”, thereby altering the ODV archive; however, the analysis presented in this paper is not significantly affected because EPA has only rarely concurred in such removal in the present study region. Section S6 of the Supplement more fully discusses this issue.

Notably, very few sites report continuous MDA8 or ODV records over the entire monitoring period, with some sites operating for only a few years. Generally, we consider all reported values in our analysis, even if only a single ODV was reported from a particular site; any temporal discontinuity associated with initiation and termination of an individual site is assumed to still allow an accurate quantification of temporal trends within the selected region. Sensitivity tests have shown that analyses based on single sites with continuous records over the entire time period agree well with analyses of records combined from multiple sites with shorter duration records within the same region.

Moved down [1]: The ozone distribution in the troposphere results from the interactions of a great many chemical and physical processes driving ozone sources, sinks and transport; fully understanding this distribution requires that the impacts of each of those processes be accurately simulated and results fully integrated with each other. Chemical transport models (CTMs) are essential to this understanding; they approach this challenging task through computational parameterizations of each process in concert with the intention of reproducing observed characteristics of the ozone distribution. However, biases in present model results are quite often found to be in the range of 5–20 ppb (Hu et al., 2017; Zhang et al., 2020). Comparison of results between different model approaches demonstrates that large uncertainties remain in our understanding; Section S1 of the Supplement briefly discusses some model result comparisons. Such comparisons have led to the increasing recognition that CTMs are not yet able to provide accurate estimates of atmospheric ozone concentrations without imposing additional constraints directly from observations. For example, Skipper et al. (2021) present a method to fuse ozone observations with USB simulated by a regional CTM in order to correct model bias, and Hosseinpour et al. (2023) present a machine learning technique that improves CTM estimates of US background ODVs, which led to improved agreement with the Parrish et al. (2017) estimate for Los Angeles and the Parrish and Ennis (2019) estimate for New York City. The much simpler, observation-based model we use takes a contrasting approach; it avoids detailed process simulation by analyzing the measured ozone distribution, which necessarily reflects the accurately integrated impacts of all relevant processes, because the atmosphere itself has performed that integration. These results also have significant uncertainty, which arises from the underlying assumptions required to interpret observations so as to extract information regarding the processes that produced the observed ozone distribution. Notably, the simplicity of this model is helpful, because impacts of the model assumptions can be readily evaluated; Sections S2–S6 of the Supplement present some of these evaluations. In contrast, CTMs require a manifold of parameterizations to simulate the many relevant chemical and physical processes; these parameterizations are generally embedded within the model computer code, rendering it extremely difficult to evaluate the uncertainty that arises from any specific parameterization. For example, evaluations by Derwent et al. (2016; 2018; 2021) identify important uncertainties in CTM results, but determining avenues to correct the model deficiencies was not possible. Neither CTMs nor our simpler approach alone can give the desired full understanding of the tropospheric ozone distribution. To advance our understanding, we believe that two key points made by Held (2005) are particularly important: “It is fair to say that we typically gain some understanding of a complex system by relating its behavior to that of other, especially simpler, systems. For sufficiently complex systems, we need a model hierarchy on which to base our understanding, describing how the dynamics change as key sources of complexity are added or subtracted.” “Elegance versus Elaboration. An elegant model is only as elaborate as it needs to be to capture the essence of a particular source of complexity, but is no more elaborate.” Emanuel (2020) makes similar points from a somewhat different perspective: “As it becomes easier to undertake complex computer simulations of climate and weather, ... it is tempting to use computers to simulate, rather than understand, nature.”

320 The ozone data considered here include the time period of emissions reductions resulting from societal efforts to control
the COVID-19 pandemic. Many publications have examined the impacts of those emission reductions on ozone in areas
throughout the world, with widely varying findings. No consistent impact has been found for summertime maximum ozone
concentrations (e.g., Gkatzelis et al., 2021), which are the focus of this study. We find no apparent systematic deviations in
MDA8 concentrations or ODVs reported for 2020 – the year of largest emission reductions – in any of the time series examined,
325 so this issue is not considered further in this analysis.

3 Methods

3.1 Quantification of temporal changes in ODVs and MDA8 distribution percentiles

Our primary analysis relies on quantifies long-term changes in ODVs recorded during the entire periods of ozone monitoring
in the southwestern US, surrounding states, and in two contrasting states in the eastern US. Since the ODVs are 3-year averages,
330 their time series are insensitive to diurnal and seasonal variations, but do reflect changes on decadal (systematic long-term
changes) and sub-decadal (i.e., interannual variability) time scales. Fits of simple, continuous functional forms to the time
series provide objective quantification of the average long-term changes. The parameter values derived from fits of Equation
3 are the basis for discussion of regional similarities and differences in ODV time series. The analysis utilized here is the
generally the same as that of Parrish et al. (2022), which is reviewed in the Introduction and Background Section 1, with the
335 additional development of an estimate for the average, long-term contribution of wildfire emissions to urban ODVs (Section
3.2). We also fit Equation 1 to the ODVs recorded at some isolated rural sites in the western US (Section 4.1) to verify that
the b and c parameter values quantified by Parrish et al. (2020) are appropriate for general application to southwestern US
ODVs.

In addition, we analyze the distribution of MDA8 ozone concentrations recorded in four coastal CA air basins; these
340 concentrations vary between days and seasonally, as well as on the longer time scales that affect the ODVs. Since our focus is
on maximum ozone concentrations, we only consider the single largest MDA8 ozone concentration observed at any of the
sites in a given air basin on each day during the ozone season, which we take as May through September. Note that the site
recording the largest MDA8 in each basin can vary from day to day. This selection gives 153 MDA8 values in each year in
each basin. From these 153 values, we characterize each year's distribution of these values by calculating seven percentiles of
345 the distribution: minimum, 10th, 25th, median, 75th, 90th, and maximum. The time series of each of these percentiles can then
be fit to the same continuous functional form (i.e., Equation 3) as fit to ODV time series. The a and A parameter values derived
in these fits then provide estimates of the year 2000 USB contribution and the US anthropogenic enhancement for the respective
percentiles. Parrish et al. (2016) utilized a similar approach in their analysis of the largest MDA8 ozone concentrations recorded
in the South Coast Air Basin (SoCAB), one of the coastal CA air basins also considered in this work.

350 Quantified uncertainties of the parameter values derived from the functional fits are important in our analysis; 95%
confidence limits are utilized, unless indicated otherwise. These confidence limits are derived from the fitting procedures

utilized in the analysis. The confidence limits from the ODV fits are widened to account for the known covariance between the recorded ODVs. Each ODV is a three-year running mean, so only every third ODV is independent from the others at a given site. Consequently, the number of independent ODVs in each fit is approximately a factor of three smaller than the number of reported ODVs, and the fitting routines therefore underestimate the true confidence limits of the derived parameters. All reported confidence limits are increased by a factor of $3^{1/2}$ to account for this covariance (Parrish et al., 2022). There may exist additional sources of covariance (regionally coherent interannual variability, temporal interannual variability, etc.) between values included in any particular fit; we cannot account for the influence of any such additional covariance.

An important issue in the present analysis is the derivation of the value of τ . The time series of urban ODVs investigated here do not allow simultaneous derivation of precise values for all three (a , A and τ) parameters of Equation 3. This difficulty is surmounted by assuming that the value of $\tau = 21.8 \pm 0.8$ years derived in earlier work for southern California (Parrish et al., 2022) is also appropriate for all regions considered here. A justification for this assumption is that precursor emission control strategies, particularly for on-road vehicles, implemented throughout the US have been similar to those in California; it is on-road vehicle emissions that have dominated urban ozone production over the past several decades (Nopmongcol et al., 2017). Parrish and Ennis (2019) also made this assumption for the northeastern US, and presented several consistency checks that show this assumption is appropriate for that region, which is further removed from southern California than the southwestern US considered here. Section S5 of the Supplement further discusses the justification for this assumption in the southwestern US and the uncertainty of the derived value of τ .

3.2 Long-term temporal changes in wildfire contributions

ODV contributions from wildfire emissions can affect the results of analysis based on Equation 3. These emissions have not been systematically reduced - rather wildfire emissions are increasing as the climate warms (e.g., Westerling et al. 2006; Westerling, 2016). Parrish et al. (2022) discuss three characteristics of the impact of wildfire emissions on ODV time series: first, wildfire emissions alone do not significantly elevate ODVs, but when they mix with local NO_x emissions, such as in central urban areas, their impact can be discerned (e.g., McKeen et al., 2002; Parrish et al., 2022; see Section 5.5 for further discussion); second, wildfire impacts have systematically increased over past decades; and third, wildfires are highly sporadic, both temporally and spatially. Addition of another term to Equation 3 can account for the systematic, temporally increasing influence of wildfire emissions. Burke et al. (2021) estimate that over the past four decades, the wildfire burned area in the US has roughly quadrupled in an approximately linear fashion (their Fig. 1A). Similarly, Iglesias et al. (2022) estimate annual mean area burned in the western US has risen by 220%–330% across a 20 year span within the period from 1984–2018 (their Fig. 7A). We represent the decadal-scale, average ODV contribution from wildfires by a similar increase; Equation 4 is Equation 3, with an added term that increases linearly by a factor of 4 from 1980 to 2020. Here the parameter WF represents the location specific ODV enhancement due to wildfires in the year 2000, and A_{WF} represents a revised A parameter in locations where enhancements of ODVs due to wildfire emissions are significant.

Deleted: the

Deleted: necessarily

Deleted: more easily

Deleted: approximately

$$\text{ODV} = a + 0.20*t - 0.018*t^2 + A_{WF}*\exp(-t/\tau) + WF*(1 + 0.03*t) \quad (4)$$

390 The linear increase of the wildfire contribution does not account for the sporadic character of wildfires; however this functional form is appropriate for the 3-year averaging period inherent in ODV time series. There are available data bases giving the spatial and temporal distributions of area burned by wildfires with detail that varies from annual means for the entire country (e.g., <https://www.nifc.gov/fire-information/statistics/wildfires>) to monthly means for individual states (e.g., monthly area burned in California from 1972 to the present is available from California Department of Forestry and Fire Protection). However, to incorporate such detailed information into our analysis would require knowledge of the spatial and temporal variability of wildfire impacts on maximum ozone concentrations at individual sampling sites, which is not available without detailed atmospheric transport modelling. Therefore, year-to-year variation of wildfire impacts undoubtedly contributes to the variability of ODVs about the functional fits to the ODV time series discussed in this work; thus, higher ODV variability in an area may be indicative of an important wildfire influence.

In principle, fitting Equation 4 to an ODV time series would allow determination of values for three parameters (a , A_{WF} and WF , with τ already fixed), but in practice such fits generally do not allow precise determination of all three parameters. However, if a can be determined from a separate analysis, then precise values can be determined for A_{WF} and WF from fits of Equation 4. We follow this approach in the analysis of time series of maximum ODVs recorded in southwestern US urban areas, which are the only areas in the region with large enough local NO_x emissions to cause significant enhancements of ODVs due to wildfire emissions. Additional complications can arise from ozone contributions produced from agricultural emissions or from other anthropogenic emissions that have not been as effectively controlled as other anthropogenic emissions. Parrish et al. (2017, 2022) discuss such ODV contributions in intense agricultural regions of California; using GEOS-Chem Geddes et al. (2022) modeled significant NO_x emissions from agricultural soils throughout the southwest but especially in Texas and California. Parrish and Ennis (2019) discuss possible ODV contributions from volatile chemical products; their emissions history is not well quantified, but are likely increasing (McDonald et al., 2018). Other ODV contributions that have not been controlled include those from oil and gas extraction processes, which are important in some regions of the southwestern US and have increased overall during the past decades; the Front Range of Colorado, including Denver (e.g., McDuffie et al., 2016) and Dallas-Fort Worth (e.g., Ahmadi and John, 2015) are urban areas where such impacts have been subject of several studies. Depending upon their specific temporal dependence, ODV contributions from these uncontrolled emission sources could contribute to the “wildfire” term in Equation 4 if they have been increasing, but none are expected to be doubling in magnitude every 20 years as the parameterized wildfire source is here. Section S4 of the Supplement further discusses some issues regarding uncontrolled anthropogenic ozone precursor emissions.

3.3 Additional analysis considerations

420 Networks of ozone monitoring sites have operated in all of the largest US urban areas; the ODVs from several of these networks are examined in detail. Where possible, the full time periods of the data records are included in the analyses. However, the analysis based on Equations 3 and 4 can only represent the time period over which ODVs consistently decreased.

In some urban areas consistent decreases did not begin until after measurements began, or the early data do not appear to accurately represent the overall urban area; in this case the ODVs selected for analysis begin at a later year. The most extreme example of the latter case is Las Vegas, where several newly established measurement sites began reporting ODVs in 2000; these values were significantly larger than reported from any other sites in that urban area. Thus, the analysis of the Las Vegas data is limited to 2000-2021 to avoid a discontinuity in the time series of ODVs. In the following section, the figures that illustrate the analyses include all recorded ODVs, so selection of ODVs for analysis can be examined.

The ODV time series fit to Equations 3 and 4 are either from single sites or multiple sites that are in reasonably close spatial proximity and exhibit similar temporal changes. There is some subjectivity in this selection of time series, which is guided by maximizing the temporal length of the series, minimizing confidence limits of the derived parameters, and minimizing deviations between the ODVs and the fits.

In addition to the confidence limits for derived parameter values, the quality of the fits of Equations 3 and 4 are quantified by root-mean-square deviations (RMSDs) between the fits and the recorded data, which scatter both above and below the fits; these RMSDs are generally in the range of 3 to 5 ppb (e.g., Tables S1 and S3-S5). These deviations are attributed to quasi-chaotic, varying ODV contributions from wildfires, stratospheric intrusions, variable meteorological conditions, etc. Our observation-based model does not account for this residual variance.

4 Results

In the following sections we analyze ODV time series recorded throughout the southwestern US and Texas and as well as in surrounding and more distant states, and the full distribution of the largest MDA8 ozone concentrations in four CA coastal air basins. Section 4.1 investigates the appropriateness of applying the b and c coefficients derived by Parrish et al. (2020) to the analysis of ODV time series recorded in the southwestern US. Section 4.2 analyzes the temporal evolution of the distribution of the MDA8 ozone concentrations in the CA air basins, and the following three sections present the analysis of ODV time series in the southwestern US (Section 4.3) Texas urban and rural areas (Section 4.4), as well as some in other US states to provide context (Section 4.5).

4.1 Long-term ODV changes at isolated rural sites in the western US

Investigation of time series of ODVs at remote rural sites provides an opportunity to quantify the long-term changes of ODVs in regions where the US anthropogenic ODV enhancements are small. In the limit of zero US anthropogenic ODV enhancement, Equation 3 is reduced to Equation 1. Here, we first fit Equation 1 to ODVs from such sites to derive b and c parameter values specific to the western US ODVs and compare those values to the values cited above from Parrish et al. (2020), and second, fit Equation 3 to these same time series to determine the small US anthropogenic ODV enhancement at those sites. Figure 1 shows ODV time series recorded at eight such sites included in the Clean Air Status and Trends Network (CASTNET) of the US EPA (<https://www.epa.gov/castnet>) whose locations are shown in inset map. They are chosen to span

Deleted: of

a wide latitude range (32° to 48.5° N), and to be as isolated as possible from anthropogenic emission sources. Seven are in a relatively narrow longitude band (109.4° to 114.2° W) located ~ 750 to 1100 km east of the US west coast, and the other is nearer (~ 220 km) the coast (Lassen Volcanic National Park (NP) at 121.6° W). All are at similar elevations (2.0 ± 0.43 km) except Glacier NP at 0.96 km.

Data from most of these same CASTNET sites have been included in previous studies of long-term ozone changes at western US rural and remote sites. Lassen Volcanic NP was investigated by Jaffe et al. (2003), Jaffe and Ray (2007), Parrish et al. (2009; 2012; 2017; 2020) and Cooper et al. (2014), due to the site's location near the US West Coast. Another four of the sites (Glacier NP, Yellowstone NP, Craters of the Moon National Monument (NM) and Canyonlands NP) were also considered by Jaffe and Ray (2007). Two of the remaining sites (Great Basin NP and Grand Canyon NP) were included by Cooper et al. (2020) in their analysis of surface ozone trends at globally distributed remote sites. Here we also consider Chiricahua NM, a more southerly site in Arizona. To our knowledge, these data sets comprise all of the longest, high quality, relatively isolated rural data sets available in the western continental US.

The CASTNET time series of ODVs are illustrated in the upper graph of Figure 1. The curves are regression fits of Equation 1, which quantify the long-term changes of baseline ozone. Table S1 gives the derived parameter values, along with confidence limits and the root-mean-square deviations (RMSD) of the ODVs from the fits. The derived a parameter values show that the absolute ODV values exhibit a systematic spatial gradient; there is close agreement among the largest a parameter values at the five southern sites (weighted mean = 71.5 ± 0.8 ppb, standard deviation = 1.1 ppb), with values decreasing with increasing latitude and/or decreasing elevation at the three more northern sites, reaching a minimum of 54.9 ± 1.1 ppb at Glacier NP, the most northern and lowest elevation site. In contrast, the derived b and c coefficients at all sites agree within their confidence limits, which indicates the absence of a statistically significant difference in the temporal evolution of the ODVs among these widely separated sites, although substantial uncertainty remains in the determination of the coefficient values.

The similarity of the temporal evolution of these rural ODVs suggests normalizing the ODVs to remove the spatial gradient and to derive a single fit of Equation 1 to all ODVs from the eight sites; the greater number of ODVs in a single fit then allows more precision in the determination of the derived parameter values. We choose to normalize to 71.4 ppb in the year 2000; this normalization factor is the average of the a parameters derived at Great Basin NP, Canyonlands NP and Grand Canyon NP (solid symbols in Figure 1), which are chosen to represent the central southwestern US. The 212 normalized ODVs recorded over 32 years at the eight sites are fit within a RMSD of 2.4 ppb (lower panel of Figure 1). This fit gives $b = 0.07 \pm 0.13$ ppb yr^{-1} and $c = -0.015 \pm 0.005$ ppb yr^{-2} ; Table S1 indicates that all of the b and c parameters derived from the separate site fits agree with these values within their confidence limits. These parameter values indicate that the rural ODVs increased early in the data record, reached a maximum in $\text{year}_{\text{max}} = 2002 \pm 4$, and decreased thereafter. A linear fit to all of the normalized ODVs over the 2000-2021 period indicates a statistically significant average negative trend equal to -0.30 ± 0.10 ppb/yr.

The b and c parameter values derived here from the normalized CASTNET ODVs agree (within derived confidence limits, see Table 1) with the parameter values derived for the entire northern midlatitudes (Parrish et al., 2020). This agreement indicates that ODVs recorded at the eight CASTNET sites provide approximate, direct measures of southwestern US

background ODVs and strongly support applying Equation 3 for analysis of ODV time series. (The derived d parameter values in Table 1 indicate that a cubic term is not justified in the polynomial fit in either the analysis of Parrish et al. (2020) or the CASTNET ODVs.) The fit of Equation 3, which includes the exponential term, to the normalized CASTNET ODVs is indistinguishable from the fit of Equation 1 (black curve in the lower graph in Figure 1), although it does give a smaller value for the year 2000 US background ODV ($a = 68.5 \pm 1.5$ ppb, compared to 71.3 ± 0.8 for the fit to Equation 1) with a small, but significant anthropogenic ODV enhancement ($A = 2.8 \pm 1.9$ ppb). We interpret the final term of Equation 3 as the mean ODV enhancement due to transport of ozone from US anthropogenic precursors to these isolated rural sites in the western US.

In summary, the 2nd and 3rd terms of Equation 3 provide an accurate estimate for the long-term change of the mean US background ODV in the western US; these terms added to the derived a parameter value of 68.5 ± 1.5 ppb, is plotted as the grey curve in the lower graph in Figure 1. Similar curves, normalized to a parameter values derived in other fits, will be included in later graphs to illustrate the purely baseline contributions to ODV time series considered in the following analyses.

4.2 Contributions to MDA8 ozone concentration distributions in four coastal California air basins

Previous analyses have shown that Equation 3 (or a closely related equation with a constant US background ODV term, rather than the small changes quantified by the 2nd and 3rd terms of Equation 3) gives excellent fits to ODV time series (i.e., fits that capture a large fraction of the ODV variance and/or with relatively small RMSDs), not only in urban areas (Parrish et al., 2017, 2022, Parrish and Ennis, 2019), but also in relatively isolated northern rural states (Parrish et al., 2022) and at western US rural CASTNET (preceding section) and coastal (Parrish et al., 2022) sites. Here we examine how well fit are the entire distributions of largest MDA8 ozone concentrations in four coastal CA air basins. These basins are selected to span a wide range of environments - the intensely photochemically active, highly urbanized South Coast Air Basin (SoCAB, containing the Los Angeles urban area) and San Diego AB, the highly urbanized but less photochemically active San Francisco Bay AB (SFB AB), and the rural North Coast AB - and to minimize impacts from uncontrolled anthropogenic emission sources, such as agricultural activity. Figure 2 illustrates the temporal evolution of seven percentiles of the MDA8 distributions; fits of Equation 3 to the time series of each percentile are included, with the derived parameter values given in Table S2. The AB locations are indicated in a map inset in Figure 3.

The fits of Equation 3 are limited to the 36 year 1980-2015 period for most of the percentiles in the three urbanized air basins; later years are not included due to clear positive deviations of the higher percentiles of the distributions, most clearly in the SoCAB, whose cause(s) have not been established. One likely cause is the rise of a new regime of wildfire impacts. According to CalFire (FRAP, <https://www.fire.ca.gov/what-we-do/fire-resource-assessment-program>) 2007 was the first time in recorded history that California lost over one million acres in a year to wildfires, but since then the four highest wildfire years have occurred in 2020, 2021, 2018, and 2017 respectively burning 4.1, 2.2, 1.8, and 1.3 million acres. Although exact matches cannot be expected without detailed analysis of where each wildfire had its impacts, these years, particularly 2020, generally stand out in the recent maximum values evident in Figure 2. In contrast, Kim et al. (2022) present a detailed investigation of these deviations in the SoCAB, and conclude that they are caused by substantial recent changes in the

520 photochemical regime in the SoCAB. Note that the fits to the lowest percentile (i.e., the minimum MDA8) in the three
urbanized air basins and to all percentiles in the rural North Coast Air Basin the fits in Figure 2 include the entire 43 year 1980-
2022 period.

Despite the marked differences in the long-term changes between percentiles and basins evident in Figure 2, the fits of
Equation 3 provide faithful descriptions of the long-term changes for all percentiles in all four basins; all RMSDs of the data
525 about the fits are in the 2.5 to 9 ppb range (Table S2), except for the maxima, which exhibit larger variability. As expected,
the anthropogenic enhancements (quantified by the derived A parameter values) are largest in the SoCAB for all percentiles,
except the minimum. The corresponding A parameter values are approximately a factor of ~ 2 smaller in the San Diego AB
than those in the SoCAB, and they are lower by another factor of ~ 2 in the SFB AB. In the North Coast AB all derived A
parameter values are near zero (-2.8 to 4.6 ppb); a significant negative A parameter value indicates that reaction of fresh NO_x
530 emissions with USB has lowered observed MDA8 concentrations below those that would result from USB alone. Importantly,
the fits to the North Coast AB percentiles are all qualitatively similar to the fits to the CASTNET ODVs in Figure 1, exhibiting
temporal behavior characteristic of baseline ozone; the fits to the minima in all four ABs also show qualitatively similar
behavior with small A parameter values ($< \sim 5$ ppb). This similarity indicates that transported baseline ozone concentrations
dominate the lowest percentiles of observed MDA8 concentrations in all air basins, even in the SoCAB.

535 The derived a and A parameter values from the fits in Figures 2 define the distributions of USB concentrations and US
anthropogenic enhancements in the year 2000; Figures 3a and 3b show bar graphs illustrating those distributions. By 2015 the
distributions of these two ozone contributions had evolved as described by Equation 3 to those shown in Figures 3c and 3d.
The US anthropogenic enhancements had decreased by a factor of 2.0 ($= \exp\{15/21.8\}$, based on the exponential term in
Equation 3), while the USB had decreased by only 1 ppb (difference in the first three terms of Equation 3 between 2000 and
540 2015). Figure 3e illustrates the mean USB distribution, i.e., the average of the five separate USB distribution estimates in
Figure 3c, which is discussed further in Section 5.1.

4.3 ODV contributions in southwestern US

The map in Figure 4 shows the ozone monitoring site locations in seven southwestern US urban areas: four larger (Phoenix
AZ, Denver CO, Las Vegas NV and Salt Lake City UT) and three smaller (Tucson AZ, Reno NV and the Albuquerque/Santa
545 Fe NM area). Monitoring sites also sparsely cover the rural areas in these five states: 19 in CO, 9 in NM, 6 in UT, 8 in AZ and
2 in NV as shown on the map in Figure S1. We examine the ODVs recorded at all of these urban and rural sites.

Figure 5 displays the analysis of the Denver CO urban (left graph) and CO rural (right graph) ODV time series. The urban
ODVs generally fall above the fit to baseline data (black curve with dashed extension, which is reproduced from the grey curve
in Figure 1). One exception is the Camp site with some ODVs falling well below the baseline curve; since this site is located
550 at street level in central urban Denver, we attribute these small ODVs to reductions in observed ozone concentrations due to
its reaction with fresh NO_x emissions. The rural data generally scatter about the baseline curve. Two rural sites that have
received attention in previous work are emphasized as solid symbols. Gothic CO in west central CO (location indicated in

Figure S1), has been considered to be a remote site (Cooper et al., 2020); that ODV time series agrees with the baseline curve within a RMSD of 1.8 ppb. Rocky Mountain NP (location also indicated in Figure S1), has been considered to be a high-
555 elevation baseline site (Lin et al., 2017). The green dashed curves show fits of Equation 3 to all ODVs in the urban (excluding the Camp site) and rural (excluding the Rocky Mountain NP site) data sets. The red curve shows the fit of Equation 4, which includes possible wildfire influences, to the maximum urban ODVs. The parameter values from these fits are annotated in the figure, and given to greater precision in Tables S3 and S4.

The derived a parameter values from analysis of Denver urban (69.0 ± 2.1 ppb) and CO rural (69.0 ± 3.1 ppb) ODVs are
560 consistent with each other and agree closely with that from the fit to the normalized CASTNET data (68.5 ± 1.5 ppb). The one exception is the Rocky Mountain NP site; the ODV temporal changes at this site are consistent with those of CASTNET data, but give a larger a parameter value (73 ± 5 ppb), which is possibly due to the relatively higher elevation of the Rocky Mountain NP site (2.7 km elevation) relative to the lower elevation of the Denver urban area (~1.8 km elevation), and/or to the influence of Denver area urban pollution, which has been demonstrated to impact this site (Evans and Helmig, 2017). The derived A
565 parameter values provide estimates of the US anthropogenic ODV enhancement in the year 2000 averaged over the sites in each group: 8.0 ± 1.7 ppb and -1.9 ± 4.4 in CO urban and rural areas, respectively; note that this rural value is consistent with zero within the derived confidence limit. The fit of Equation 4 to the time series of maximum ODVs (with the a parameter value fixed at 69.0 ppb) gives a somewhat larger A parameter value ($A_{WF} = 11.0 \pm 1.7$) and a WF parameter value of 4.0 ± 2.5 ppb; this latter value is an estimate of the ODV enhancement due to wildfire emissions in the year 2000.

An open scientific question is the role of emissions from the oil and natural gas industry and agricultural activities in
570 elevating ODVs in the Denver urban area. To the extent that emissions from these sectors have increased similarly to wildfires, any such role would contribute to our derived WF parameter value, which is larger in the Denver urban area than other urban areas considered here (see Table S4). With a different temporal dependence they could bias the derived a parameter value; however, the good agreement between that estimate in the Denver urban area, rural CO and the CASTNET data indicates no
575 more than a small bias.

Figure 6 shows the results of similar analyses for six other southwestern US urban areas. In these plots (and in Figure S2),
the black, generally lower curves with dashed extensions are the fit to the baseline data from Figure 1, normalized to the
respective a parameter values; thus, these curves indicate the temporal behavior of ODVs in each data set that would have
been expected in the absence of US anthropogenic emissions. The derived a parameter values are similar in 5 of those areas,
580 all falling in the range of 66.2 to 69.0 ppb, and agree well with the Denver (69.0 ± 2.1 ppb) and CASTNET (68.5 ± 1.5 ppb) results discussed above. Only in Tucson is a significantly smaller a parameter value (63.9 ± 1.4 ppb) derived; this value is also smaller than found at the nearby Chiricahua NM CASTNET site (70.1 ± 1.7 ppb); the lower elevation of Tucson may contribute to this difference. Fits of equation 3 to all of the rural ODV time series (Figure S2 and Table S3) also give similar a parameter values (within 65 to 69 ppb), with small A parameter values (-3 to +5 ppb). Fits of equation 4 to the maximum urban ODVs
585 (with the a parameter values fixed at the values derived from the ODVs in the entire respective urban area) give A_{WF} parameter

values in the range of 6 to 16 ppb. Small, positive wildfire ODV enhancements (WF = range 0.6 to 1.6 ppb) are derived in all six urban areas, although the confidence limits of all include zero.

In summary, analyses of the ODV time series throughout the southwestern US give remarkably consistent results. The weighted average a parameter value, which gives an estimate of the US background ODV in the year 2000, derived from 11 urban and rural ODV time series, is 67.4 ± 0.7 ppb with only the low elevation, most southerly Tucson urban area giving a significantly lower value; this average agrees well with the value derived in the CASTNET site analysis (68.5 ± 1.5 ppb). This overall agreement indicates that an approximately constant, common US background ODV value is found throughout the southwestern US. The mean A parameter value, which gives an estimate of the US anthropogenic ODV enhancement, derived from four rural ODV time series, is -0.1 ± 2.9 ppb, which indicates that ODVs outside of urban areas are little affected by US anthropogenic emissions throughout this entire region. Any area with intense agricultural activity may be an exception to this general finding, due to the influence of uncontrolled anthropogenic emissions from this activity. Within the seven urban areas, the A_{WF} values span the range from 6 to 16 ppb, with the larger values (11 to 16 ppb) derived in the four nonattainment areas. Wildfire emissions contribute additional ODV enhancements within urban areas: estimated as 0.6 to 4.0 ppb (mean of 1.6 ± 0.9 ppb) in the year 2000, and 1 to 6.4 ppb (mean of 2.6 ± 1.4 ppb) in 2020.

600 4.4 ODV contributions in Texas

The ozone monitoring site locations throughout Texas, plus some sites in neighboring states, are indicated in Figure 7, which also identifies the division of these sites among nine Texas regions and the state of Oklahoma; we examine the ODVs recorded in these regions. Figure 8 plots the ODV time series recorded in seven of the nine Texas regions, and Figures S3 and S4 give similar plots for the other two Texas regions, for Oklahoma, and for eight additional surrounding states. The parameter values derived from fits of Equations 3 and 4 to all ODV time series are annotated in the figures, and given to greater precision in Tables S4 and S5. In Figure 8 and in the upper graphs of Figure S3 the ODVs for the entire state of Texas (grey symbols) are included in the plots highlighting the regional ODVs (colored symbols), which are fit to Equation 3 as indicated by the upper black solid curves. Lower black curves with dashed extensions indicate the fit to the baseline data from Figure 1, normalized to the respective a parameter values; these curves indicate the temporal behavior of ODVs in each region that would have been expected in the absence of US anthropogenic emissions.

Comparison between Figures 6 and 8 (note differences in ordinate scale) indicates that ODVs in Texas were often much larger in earlier decades than those in the southwestern US urban areas, indicating much larger impacts from US anthropogenic emissions in Texas. For example, in 2000 the maximum US anthropogenic ODV enhancements (i.e., A_{WF} parameter values) in Texan urban areas were generally > 25 ppb, and as much as 54 ± 3 ppb in the Houston region, compared to the range of 6 to 16 ppb found in the Southwestern US. However, the ODVs in Texas have decreased by a larger absolute amount than in the southwestern US, so that at present maximum ODVs in Texas are no larger than those in the Denver and Phoenix urban areas.

The southwestern US is characterized by an approximately constant US background ODV across the entire region, but across Texas there is a pronounced spatial gradient. The a parameter values of 65 ppb derived for the El Paso and Western

620 Rural regions are similar to the weighted mean (67.4 ± 0.7 ppb) of the urban and rural southwestern a parameter. However, the US background ODV decreases with distance both east and south across the state, with the smallest values found in southern and eastern Texas - 50 ± 5 ppb in SW Texas and 54 ± 3 ppb Houston - which is heavily influenced by transport of maritime air from the Gulf of Mexico (Berlin et al., 2013) during the warm (ozone) season. This broad geographic feature in USB is also seen in several modelling studies (e.g., Fiore et al., 2014; Zhang et al., 2020).

625 As expected, the largest Texas US anthropogenic ODV enhancements (27 to 54 ppb in 2000) have generally been found in the nonattainment areas within the state - Houston, Dallas and San Antonio – although comparably large A parameter values (28 to 37 ppb) are found throughout southeast and east Texas. It is notable that the El Paso region is anomalous – although the A parameter value (11 ± 2 ppb) is comparatively modest, the a parameter value (65 ± 2 ppb) is so large that even the relatively small US anthropogenic ODV enhancement is sufficient to produce ODVs that routinely exceed the 70 ppb NAAQS throughout the entire measurement record (upper left graph in Figure 8).

630 There is evidence that the maximum ODVs in Texas urban areas may be elevated by ozone sources that have not yet been effectively controlled by air quality improvement efforts. Such sources could possibly include wildfire emissions, which are approximately quantified by Equation 4, or emissions from oil and gas exploration and production, which are ubiquitous across Texas. To gauge the possible magnitude of these sources, Equation 4 is fit to the time series of maximum ODVs in Dallas and Houston (dotted black curves in the middle graphs of Figure 8). These fits possibly indicate contributions of 2 to 3 ppb to the maximum ODVs in these urban areas from such growing, uncontrolled anthropogenic sources. Iglesias et al. (2022) show the significant growth in wildfires across all of Texas (their Figure 2), especially since the turn of the century, and Gong et al. (2017) calculate that wildfires impact Houston on 3.5% of the days in their May-September 2008-2015 study period, with an average contribution of ~ 8.5 ppb.

4.5 ODV contributions in neighboring and more distant states

640 For context, preliminary analyses have been conducted of the ODV time series recorded in states lying east and north of the region that is the focus of this study. Figures S3 and S4 illustrate the results of these analysis for Oklahoma, Louisiana, Arkansas, Nebraska and Kansas; Equation 3 is fit to the time series of all ODVs recorded in each state without consideration of intrastate regional differences. Nevertheless, the RMSD of the ODVs about the fits are similar to the fits to more carefully selected sets of sites discussed in the preceding sections. The results for these states are considered to be adequate for determining boundary conditions for the ODV component determinations in the region that is the focus of this work.

645 Figure S3 also includes analyses of ODVs recorded in four rural northern states: Montana, North and South Dakota, and Wyoming. A fit to all ODVs recorded in these four states gives a mean A parameter value of 1.3 ± 1.7 ppb, which indicates only small US anthropogenic ODV enhancements throughout that region. (Note: the Boulder site in Wyoming – AQS Site ID 56-035-0099 reports anomalously large ODVs; since this site is in the Upper Green River Basin where high ozone concentrations are produced in wintertime from the region's large oil and gas production activities (Schnell et al., 2009), all ODVs from this site have been excluded.) Separate fits of Equation 3 to the ODV values in each state provide separate a

parameter values; there are some small differences in these results: a maximum in Wyoming of 64 ppb, a minimum in Montana (59 ppb), with North and South Dakota being intermediate (60 and 62 ppb, respectively). Parrish and Ennis (2019) and Parrish et al. (2022) present earlier analyses for three of these states, and found results consistent with this updated analysis.

655 Figure S6 includes analyses using fits of Equation 4 to the time series of maximum ODVs recorded in the New York City and Atlanta urban areas. The New York City time series was originally analyzed by Parrish and Ennis (2019) (see their Figure S4); here that time series is extended through 2021. Figure S6 includes separate symbols for the Atlanta, coastal and inland sites.

5 Discussion and Conclusions

660 5.1 Background ozone dominates observed concentrations throughout the US

Five estimates of the May to September USB distribution derived from analysis of observations from CA West Coast locations are compared in Figure 3c. The first bar illustrates the ozone distribution measured between 0.6 and 1.0 km altitude by sondes launched from a coastal site at Trinidad Head (Oltmans et al. 2008), which is located in the Northern Coast AB shown in map in Figure 3; Parrish et al. (2022) fully describe this data set. The ozone altitude gradient (see Figure 1 of Parrish et al., 2022) indicates that this altitude range includes the MBL to free troposphere transition, so that it is high enough to avoid continental influences as air is transported ashore and low enough to well-characterize air mixed into the convective boundary layer over the continent (Parrish et al., 2010). Thus, these sonde data provide direct estimates of USB concentrations. The other four bars show estimates of the distribution of USB ozone derived in Section 4.2 from the MDA8 distributions in CA coastal air basins. There is reasonable agreement between the five estimates, but with some notable differences. The two southern estimates (San Diego AB and SoCAB) are somewhat larger than the three more northerly, which is consistent with the systematic decrease of USB with latitude quantified by Parrish et al. (2022). Variation in site elevations also contributes to these differences. The Crestline site at 1.4 km and the Alpine site at the 0.6 km are included in the SoCAB and San Diego AB, respectively; they are at higher elevations than other coastal air basin sites, and very often record the largest MDA8 ozone concentrations in their respective AB. Thus, at least some of the quantitative differences in the bars in Figure 3c represent real atmospheric differences. The mean of all five USB distributions from that graph is included in Figure 3e; it provides an estimate for the average USB distribution along the CA coast.

Comparison of the distributions in Figures 3c, 3d and 3e shows that by 2015 the distribution of USB was larger than the distribution of the US anthropogenic ozone enhancements in all four CA coastal air basins. This dominance of background ozone is quite pronounced; only in the SoCAB and San Diego AB do the distributions in Figure 3d overlap at all with the mean USB distribution in Figure 3e; in the SoCAB the 75th percentile of the US anthropogenic ozone enhancement is smaller than the 10th percentile of the mean USB contribution. Comparisons for the separately derived air basin USB distributions between Figures 3c and 3d shows that the USB dominance in the SoCAB and San Diego AB comparisons are even more pronounced. This predominance of the background contribution developed relatively recently; Figure 3a and 3b show the comparison for

Deleted: an

Deleted: analysis

the year 2000, when the US anthropogenic enhancements were generally greater than or equivalent to the USB contributions in the SoCAB, and more nearly comparable in the San Diego AB.

690 The predominant contribution of USB to maximum ozone concentrations, even in the SoCAB where the largest US ozone concentrations occur, emphasizes the great difficulty of achieving accurate air quality modeling. Traditionally, CTMs have primarily focused on careful, detailed treatment of transport and photochemical processes on local to regional scales, which is required for the accurate simulation of the US anthropogenic ozone enhancements; however, less attention has been paid to the USB contributions, especially the day-to-day variability. Of great importance for present day exceedance event modeling or daily air quality forecasts, is the accurate simulation of USB on short-term, i.e., daily or even hourly, time scales. Since USB is driven by a manifold of processes occurring on small spatial scales, but over hemisphere-wide distances, accurate, 695 detailed simulation of the stratospheric and tropospheric contributions to USB is beyond the capabilities of current air quality modeling, as it is of modeling for predictions of future baseline ozone concentrations. These issues are explored in two recent comparisons of estimates of background ozone and their impacts on the simulation of surface ozone across the continental US (Hogrefe et al., 2018; Zhang et al., 2020) where background concentrations were shown to differ by 5-10 ppb in the mean and up to 15 ppb for domain-averaged MDA8 values. Figure 3e illustrates the differences between two different model simulations 700 of the US background MDA8 ozone distributions for April - June 2017 at 12 high elevation sites in the western US. As Zhang et al. (2020) note, the means of the distributions agree within 6 ppb. However, the larger percentiles of the distribution differ more widely; e.g., the 98th percentiles are ~70 ppb and ~57 ppb from the GFDL AM4 and GEOS Chem models, respectively. It's notable that the former model indicates that USB alone can exceed the 2015 NAAQS of 70 ppb, while the latter model finds that USB always remained significantly below the NAAQS.

705 5.2 Pronounced shift in spatial distribution of maximum US ozone concentrations

Great efforts have been made over the past half century to reduce anthropogenic precursor emissions throughout the entire country. These efforts have been remarkably successful, yielding substantial reductions in the number and magnitude of US ozone NAAQS exceedances; however, progress has not been spatially uniform over the country. Figure 9 illustrates one measure of this progress. During the 5-year period of 1997-2001 the large majority (43) of the 51 US states plus the District of Columbia recorded at least one ODV of 75 ppb or larger; 20 years later, during the 2017-2021 period, the number of such 710 states had been reduced to 20. In the earlier period 80% of ODVs across the Nation were ≥ 75 ppb, and in the later period that percentage had dropped to 10%, clearly demonstrating a remarkable improvement in ozone air quality. However, there are marked regional differences in the percentages of ODVs ≥ 75 ppb - 88% and 64% in the eastern and western regions, respectively, in the earlier period, but 2.4% and 22%, respectively in the later period. In Figure 9 the solid blue bars in the 715 western region demonstrate that nearly all of the seven total states that recorded more than 7% of ODVs ≥ 75 ppb were in the southwestern US, including Texas and California; the single exception is Connecticut, where the maximum ODVs are recorded near the coast of Long Island Sound (Parrish and Ennis, 2019). Similar conclusions are reached if ODVs > 70 ppb are considered (Figure S5). In summary, great improvement in US ozone concentrations have occurred over past decades, but that

improvement has been accompanied by a marked shift of the highest observed ozone concentrations from the eastern to the southwestern US.

The occurrence of only modest ODV decreases in the southwestern US has been noted in other analyses. Simon et al. (2015) found that the upper end of the summertime urban ozone distribution (i.e., representative of ODVs) generally decreased over the 1998-2013 period throughout the US in response of decreasing emissions of NO_x and volatile organic compounds (VOC). They interpreted those results as demonstrating the large scale success of US control strategies. However, their Figure S12 shows that the Denver urban area was an exception to this generality, with all sites exhibiting either positive or insignificant trends. Similarly, Abeleira and Farmer (2017) found that ozone in the Denver urban area was either stagnant or increasing between 2000 and 2015, despite substantial reductions in NO_x emissions. In Colorado Springs, approximately 100 km south of Denver, Flynn et al. (2021) found that summertime MDA8 ozone shows no significant trend throughout its distribution over the past 20 years. Further, they found little evidence of local photochemical production or ozone transport from Denver, consistent with our classification of this area as rural. The results of these three studies are consistent with the long-term ODV changes in urban and rural Colorado shown in Figure 5, and our conclusion that the US background ODV is the dominant contributor to ODVs.

5.3 Spatial and temporal distribution of ODV contributions

The derived spatial distributions of the two major ODV components –the US background ODV, and the US anthropogenic ODV enhancement – are illustrated for the year 2000 as contour maps in Figures 10 and 11. In Figure 10 the estimated US background ODVs are at a maximum - greater than 64 ppb (see violet contour) - throughout the entire southwestern US, including western Texas and Wyoming, and above 68 ppb in some areas, including most of Colorado. We attribute these large values to baseline air at higher altitudes flowing over the inland mountain ranges, and mixing into the convective boundary layer over the continent (Langford, et al., 2017). The overall spatial pattern is similar to those simulated for surface impacts of ozone transport from Asia (see Fig. 9 of Lin et al., 2012a), stratospheric intrusions (see Fig. 11 of Lin et al., 2012b) and USB distributions (see Fig. 1 of Zhang et al., 2020). Previous studies of reanalysis data sets (Sprenger and Wernli, 2003; Škerlak et al., 2014) describe the southwestern US as prone to high rates of deep stratosphere to troposphere transport. These events tend to cross the tropopause at high latitudes (40-60°N) near the end of the North Pacific storm track but make their surface impacts felt most acutely after quasi-isentropic subsiding transport to the southeast (in the lee of the Pacific High). While the peak of this effect occurs in the spring it does continue into the summer (Škerlak et al., 2014), when it can more strongly couple with the surface because of the very deep convective boundary layers that develop in the arid, high terrain of the southwestern US. With such large US background ODV values, even relatively small anthropogenic enhancements in the regional urban areas raise ODVs above the 70 ppb ozone NAAQS, thereby leading to their designation as nonattainment areas. Were the NAAQS to be lowered further, for example, to 65 ppb as has been considered in the past (e.g. [Regulatory Impact Analysis of the Final Revisions to the National Ambient Air Quality Standards for Ground-Level Ozone](#)), Figure 10 indicates that NAAQS achievement would be precluded over the vast southwestern US region inside the 64 ppb contour (unless regulatory efforts

could reduce the US background ODV). A striking feature of the contour map in Figure 10 is the significant spatial gradient in the US background ODV over the continent. Increases occur inland from the West Coast as surface elevations increase (Parrish et al. 2022), with decreases to the north and east of the broad maximum in the southwestern US, and to the east and south within Texas. These smaller values in eastern Texas (e.g., 54 ± 3 ppb in Houston) allow for larger anthropogenic ODV enhancements there before the NAAQS is exceeded.

CTMs have been extensively utilized to estimate US background ODVs. Figure 10 compares our observation-derived results with a portion of Figure 3 of Jaffe et al. (2018), which presents the results of a simulation by the GFDL-AM3 model of a statistic comparable to the US background ODV. The broad spatial features are similar in both plots; the maximum in the southwestern US is the dominant feature in each. The maximum observation-derived values (68-70 ppb) are consistent with the model simulated maximum (60-70 ppb). All of the observation-derived values, except for a narrow band along the Pacific Coast, are in the 50 to 70 ppb range, as are the model derived values, with the exception of the values in the southeast corner of that map. Both results show general south-to-north and west-to-east decreases of inland US background ODVs. However, it is apparent that the observational-based estimate has substantially greater precision and spatial resolution than provided by the low resolution (200×200 km²) model estimate. Jaffe et al. (2018) estimate that the model uncertainty of the seasonal mean USB is $\sim \pm 10$ ppb, but larger for individual days, such as is required for determination of US background ODVs. The magnitudes of the model and observational-derived results in Figure 10 agree within this estimated uncertainty of the model simulations. The US background ODV estimated from the GFDL-AM4 simulated ozone distribution (approximated by the 98th percentile) shown in Figure 3 is also consistent with the contour map in Figure 10, indicating that US background ODVs above 68 ppb over the southwestern US are indeed physically realistic.

The NO₂ column measurements from the OMI satellite (Lu et al., 2015) are included in Figure 11 for comparison with the US anthropogenic ODV enhancements, which outside CA, are generally elevated in the larger southwestern US and Texas urban areas. There is general correlation of the NO₂ columns within these urban areas, which supports our interpretation of the derived *A* parameter value as quantifying the US anthropogenic ODV enhancement. However, some areas of clear differences between those ODV enhancements and the OMI NO₂ column measurements are evident in Figure 11. Most clearly, emissions from the large coal-fired power plants in the Four Corners region (location identified in Figure S1) give a strong elevation in the NO₂ column that is not reflected in US anthropogenic ODV enhancements. This lack of correlation is consistent with the ODV time series for this region (Figure S2), which indicates that the US anthropogenic ODV enhancement is near zero. Mesa Verde NP is in this region; it has been considered a high-elevation site representing baseline O₃, at least in the spring, by Lin et al. (2017). This site records ODVs generally consistent with all other sites in this region. The co-location of the large NO₂ enhancement with an indiscernible US anthropogenic ODV enhancement indicates that the large NO₂ emissions from the power plants have only a negligible effect on the regional ODVs. The Denver urban area, with *A* parameter values in the 8 to 11 ppb range (Figure 5), does not stand out in Figure 11, although that area does have a clear signature in the OMI NO₂ column data.

785 Figure 12 compares the long-term changes of the ODV components between the Crestline site in the Los Angeles urban
area with those in the four larger southwestern US urban areas and three Texas urban areas. Since ~1990, the Crestline site has
usually recorded the largest ODV in the SoCAB. Overall, decreases in ODVs have been recorded in each of the eight cities;
this decrease is primarily driven by a decrease in the US anthropogenic ODV enhancement. The final term in Equation 3
indicates that this contribution decreased by a factor of 6.3 between 1980 and 2020. In contrast, the first 3 terms of Equation
790 3 indicate that the US background ODV increased over the entire western US region by ~11 ppb from 1980 to the mid-2000s,
and has since slowly decreased. The relatively minor, but increasing (up to ~6 ppb in Denver in the year 2020) mean ODV
enhancements due to wildfire emissions are separately indicated. Two conclusions emerge from Figure 12. First, throughout
the monitoring record, the US background ODV has been, by far, the largest ODV contribution in the four southwestern US
urban areas, and that by 2000 it had become the predominant ODV contribution in all cities, even within the Los Angeles
795 urban area exemplified by the Crestline site. This conclusion is consistent with the discussion in Section 5.1. Second, the
increase in the US background ODV between 1980 and the mid-2000s and the increasing wildfire contribution through the
entire period, partially offset the reduction in the US anthropogenic ODV enhancement, so that ODVs have not changed greatly
in the southwestern US urban areas and El Paso, Texas.

There are some notable deviations from the overall picture described above. First, the results shown in Figure 12 and all
800 fitted curves in all figures represent the average behavior of the fitted ODVs; there is significant scatter about those fits as
quantified by the RMSDs (generally < 5 ppb) annotated in the figures and given in Tables 1, S1 and S3-S5; this scatter appears
to be predominately year-to-year variability. However, one systematic deviation is particularly relevant - the recent anomaly
of the El Paso nonattainment area and nearby rural New Mexico (see graphs in Figures 8 and S3); the ODVs in the last few
years are as much as 15 ppb larger than expected from the fits to Equation 3. It is notable that in the most recent year included
805 in this analysis (2021) the largest ODV (80 ppb) in the two state region of Texas and New Mexico was not recorded in either
of the traditional urban ozone hot spots of Houston or Dallas; rather it was recorded at the rural Desert View site in New
Mexico, which is included in the El Paso region in Figure 8. Only a single site in Phoenix within the entire southwestern US
and Texas region (excluding California) equaled that ODV. ODVs nearly as high were recorded in rural New Mexico sites
that are included in the Western Rural TX region in Figure S2. The cause of these high ODVs is presently unexplained,
810 although it may be important to note that these regions are in the Permian oil and gas basin (Karle et al., 2021).

5.4 Implications for US air quality policies

This spatial difference in the success of air quality policies aimed at ODV reduction has resulted in a pronounced regional
shift of the occurrence of ODVs of 75 ppb and above (and above 70 ppb – Figure S6), with a growing preponderance of the
nation's largest ODVs recorded in the southwestern US (see discussion in Section 5.2). The cause of this change is clear in
815 Figure 10: the southwestern US suffers from large US background ODVs that closely approach the NAAQS. In Figure 12, the
US anthropogenic ODV enhancements in the southwestern US urban areas and El Paso Texas have been reduced in magnitude
to such an extent that by 2020 all were smaller than 6 ppb. However, even these remaining enhancements are large enough to

raise the recorded ODVs above the 70 ppb 2015 NAAQS, as well as the 75 ppb 2008 NAAQS. This conclusion is further illustrated in Figure 13, where fits of Equation 4 to time series of maximum ODVs in three southwestern US urban areas are compared to similar fits in three of the largest US urban areas; additionally, the fit of Equation 3 to the time series of CASTNET ODVs and the temporal evolution of the corresponding baseline ozone concentrations from Figure 1 are included. In Atlanta and New York City, with substantially lower year 2000 maximum US background ODVs of 49 and 52 ppb, respectively, the fits to the urban maximum ODVs dropped below the Phoenix and Denver fits in the early 2010s; they have now dropped below the 70 ppb NAAQS and are approaching the Reno, CASTNET and baseline fits. Interestingly, by 2021, the fit to the maximum ODVs recorded at all locations in Figure 13, from the isolated rural CASTNET sites in the western US to the major metropolitan areas of Atlanta and New York City (excluding the Los Angeles) were in the 66 to 76 ppb range, with the southwestern US urban areas defining the upper limit. Importantly, all curves included in Figure 13 are direct fits to measured concentrations; they do not depend upon accurate differentiation between the separate ODV contributions.

The spatial shift of the nation's highest ozone concentrations to the southwestern US emphasizes a long-standing concern with US policies for improving ozone air quality – a single standard is applied to the entire country without regard to the USB; this background cannot be addressed by local or regional reductions in ozone precursors. This policy contrasts to those addressing another environmental hazard – exposure to ionizing radiation. The U.S. Nuclear Regulatory Commission (NRC, <https://www.nrc.gov/about-nrc/radiation/around-us/doses-daily-lives.html>) estimates that about half of the average American radiation dose is due to natural background radiation that, similar to ozone, is higher in the high altitude southwestern US. Health and safety standards for ionizing radiation are based on the additional exposure that comes from anthropogenic sources of radiation, not on the overall radiation exposure. A similar approach for ozone would require that the ozone standard be based on the anthropogenic increment of ozone above the USB within a region, rather than on the total ambient ozone concentration; alternatively, a regionally varying ambient concentration standard that accounted for the regionally varying USB could serve that purpose. Of course, such an approach would bring additional complexity, since ozone is secondary air pollutant that can cross state lines, making it difficult to define a suitable boundary for any particular region, such as the southwestern US. However, as of 2020, we estimate that the US anthropogenic ODV enhancement in eight southwestern US urban areas (seven in Figure 4, plus El Paso TX) had been reduced to the range of 2.4 to 6.4 ppb; yet the US background ODVs, plus the relatively small wildfire contributions, are so large that 5 of these 8 cities are the centers of ozone nonattainment areas. Importantly, it is only the small US anthropogenic ODV enhancement that can possibly be directly reduced through further controls on urban and industrial emissions, although in addition the small wildfire contributions possibly can also be reduced indirectly through decreased NO_x emissions in VOC-poor urban areas. In Denver in 2020, we estimate that the US anthropogenic ODV enhancement had been reduced to 4.4 ppb, yet the US EPA has recently downgraded the Denver urban area from a "Serious" to "Severe-15" nonattainment area under the 2008 ozone NAAQS (<https://www.govinfo.gov/content/pkg/FR-2022-10-07/pdf/2022-20458.pdf>). This redesignation will require further reductions in local and regional precursor emissions even though the US anthropogenic ODV enhancement is already so small that such additional emission control efforts can be expected to have little impact on future ODVs.

Importantly, it is estimated that between 1950 and 2000 baseline ozone concentrations at northern midlatitudes increased by a factor of 2.1 ± 0.2 (Parrish et al., 2021b). These baseline concentrations, which largely account for the US background ODV, reached a maximum in the mid-2000s, and since have slowly decreased at a mean rate of ~ 1 ppb decade⁻¹ (Parrish et al., 2020). This decrease results from decreasing anthropogenic ozone precursor emissions throughout northern mid-latitudes due to implementation of effective emission controls in North America, Europe and, more recently, east Asia. Given the large increase in baseline ozone that occurred as anthropogenic emissions increased in the 20th century, it can be expected that a significant decrease in baseline ozone can be achieved by further reducing total zonal precursor emissions. Cooperative, international emission control efforts aimed at continuing, or even accelerating, the current decrease of baseline ozone concentrations, and thereby the US background ODV, may be an effective alternative policy approach to further reducing US ODVs. It should be noted that this expectation relies on the assumption that the background will continue to decrease; however that expectation could be compromised by the changing climate. For example, stratosphere-troposphere exchange rates could increase due to acceleration of the Brewer-Dobson circulation; modeling work by Abalos et al. (2020) points toward a 10-15% rise in the stratosphere-troposphere O₃ source by the end of this century.

We should step back from comparison of air quality policy approaches to note that there is evidence that further reductions of ambient ozone concentrations throughout the US will provide significant health benefits (e.g., Zhang et al., 2019). This applies to both attainment and nonattainment areas, and to the entire concentration distribution of ozone, not just the few days of highest concentrations that determine the ODVs that are the focus of NAAQS attainment. Thus, all reductions of ozone precursor emissions will be beneficial, and if carried out in all northern midlatitude countries, will serve to further reduce baseline ozone concentrations, thereby easing the attainment of air quality standards based on ODVs or other statistics designed to quantify ozone concentrations of concern. Moreover, control measures that reduce VOC and NOx emissions have the added benefits of reducing other criteria air pollutants, air toxics, co-emitted greenhouse gases, and precursors to fine particle formation, which contribute to haze and visibility reduction; further air quality improvement is desirable from multiple perspectives.

5.5 A required modelling hierarchy

Derwent et al. (2023) emphasize the need for a widely-accepted, simple, conceptual 'model' that intuitively explains the broad features of how ozone sources, sinks and transport processes all interact to establish the observed local, regional and larger-scale spatial distributions, seasonal cycles and long-term temporal changes of ozone. Broadly speaking, this conceptual model envisages background air containing ozone and other baseline trace species entering an urban area or rural region. Urban, other anthropogenic, and biogenic precursor emissions drive local and regional scale photochemical ozone production, which elevates ozone concentrations above the global baseline levels, leading to human health effects and crop and vegetation damage. After one to several days travel downwind, the regionally-polluted air with elevated levels of ozone, other photochemical products, and unreacted precursors is lofted from the continental boundary layer and rejoins the global

Deleted: Finally, w

885 circulation. Neither the conceptual ‘model’ nor our observation-based model applied in this paper can alone provide a
quantitative understanding of tropospheric ozone; rather, a hierarchy of models is required.

In this regard, we believe that two key points made by Held (2005) are particularly important:

- 890 • “It is fair to say that we typically gain some understanding of a complex system by relating its behavior to that of
other, especially simpler, systems. For sufficiently complex systems, we need a model hierarchy on which to base
our understanding, describing how the dynamics change as key sources of complexity are added or subtracted.”
- “Elegance versus Elaboration. An elegant model is only as elaborate as it needs to be to capture the essence of a
particular source of complexity, but is no more elaborate.”

Emanuel (2020) makes similar points from a somewhat different perspective:

- 895 • “As it becomes easier to undertake complex computer simulations of climate and weather, ... it is tempting to
use computers to simulate, rather than understand, nature.”
- “It is sometimes perceived as easier to run the model than to develop a comprehensive and satisfying
understanding of the phenomena in question.”

In addition to the conceptual ‘model’ and the simple mathematical model that we apply here, comprehensive numerical models
(i.e., CTMs) that aim to simulate in full detail as much of the atmospheric chemistry and dynamics as possible are essential for
900 providing the beautifully detailed results that are presently available to the atmospheric community. CTMs require great
“elaboration” to adequately treat the manifold of important processes, while our observation-based model is focused much
more on “elegance”, but it does indeed “capture the essence of a particular source of complexity”. In the present work that
complexity is the interaction of USB with US anthropogenic contributions to establish observed ozone concentrations. The
simplicity of our model allows direct, conceptual understanding of the phenomena that underlie the observed ozone
905 distribution, and this understanding provides context for the detailed results provided by CTM simulations.

From the “elaborate” perspective, it is well-understood that the temporal and spatial distribution of ozone in the
troposphere results from the interactions of a great many chemical and physical processes driving ozone sources, sinks and
transport; fully understanding this distribution requires that the impacts of each of those processes be accurately simulated
and results fully integrated with each other. CTMs approach this challenging task through computational parameterizations
910 of each process in concert with the intention of reproducing observed characteristics of the ozone distribution. However,
biases in present CTM results are quite often found to be in the range of 5-20 ppb (Hu et al., 2017; Zhang et al., 2020).
Comparison of results between different models demonstrates that large uncertainties remain in our understanding; Section
S1 of the Supplement briefly discusses some model result comparisons. Such comparisons have led to the increasing
recognition that CTMs are not yet able to provide accurate estimates of atmospheric ozone concentrations without imposing
915 additional constraints directly from observations. For example, Skipper et al. (2021) present a method to fuse ozone
observations with USB simulated by a regional CTM in order to correct model bias, and Hosseinpour et al. (2023) present a
machine learning technique that gives more accurate CTM estimates of US background ODVs; this approach led to
improved agreement with the Parrish et al. (2017) estimate for Los Angeles and the Parrish and Ennis (2019) estimate for

920 New York City. Our much simpler, observation-based model takes a contrasting approach; it avoids detailed process
simulation by analyzing the measured ozone distribution, which necessarily reflects the accurately integrated impacts of all
relevant processes, because the atmosphere itself has performed that integration. Results of our simpler model also have
significant uncertainty, which arises from the underlying assumptions required to interpret observations so as to extract
information regarding the processes that produced the observed ozone distribution. In this regard, the simplicity of this
925 model is helpful, because impacts of the model assumptions can be readily evaluated; Sections S2-S6 of the Supplement
discuss some of these evaluations. In contrast, the manifold of parameterizations that CTMs require to simulate the many
relevant chemical and physical processes are generally embedded within the model computer code, rendering it extremely
difficult to evaluate the uncertainty that arises from any specific parameterization. For example, evaluations by Derwent et
al. (2016; 2018; 2021) identify important uncertainties in CTM results, but determining avenues to correct the model
deficiencies was not possible. Neither CTMs nor our simpler approach alone can give the desired full understanding of the
930 tropospheric ozone distribution – hence the need for a robust modelling hierarchy.

Continuing to guide US ozone air quality policy development requires this modelling hierarchy to address new challenges.
For more than five decades, those policies have primarily focused on urban ozone produced from precursors emitted by mobile
and industrial sources; the contributions from transported background ozone and photochemical production driven by other
emission sources received much less attention. However, as we demonstrate in this and earlier papers, that original
935 anthropogenic contribution has been so effectively reduced (a decrease of more than a factor of six from 1980-2020), that the
background contribution now predominates throughout the US. Accordingly, increased attention must now be placed on a
variety of additional anthropogenic sources, notably including emissions from wildfires, oil and gas production, and
agricultural activities. Present-day CTMs simulate ozone production from these additional sources, but the results have
substantial uncertainties arising from the necessity of developing and incorporating a manifold of additional parameterizations
940 for their treatment. Observational-based modelling also can give insights into the influences of these additional sources without
the need for such parameterizations; these insights arise from temporal and spatial correlations between observed ozone
concentrations and emissions from these sources. For example, in this work, we quantify small but significant ODV
contributions from wildfires in central urban areas of the southwestern US; similar analyses for urban areas of the Pacific
Northwest are given by Parrish et al. (2022). In previous papers (Parrish et al. 2017; 2022) we have identified indications that
945 soil emissions of nitrogen oxides in California’s regions of intense, fertilized agricultural activity (i.e., the Imperial, San
Joaquin and Salinas Valleys) increase ODVs by 5-15 ppb. We conclude that a robust understanding of the spatial and temporal
distribution of US ozone concentrations requires a model hierarchy that includes both CTMs and observational-based models,
plus hopefully other modeling approaches. We can have confidence in that understanding only when all the models can
converge to a consistent quantification of that distribution, and only then can reliable policy guidance be provided.

Formatted: Font color: Black, English (UK)

950 **Acknowledgments, Samples, Data and Legal Notice**

D.D. Parrish is an independent consultant (David D. Parrish, LLC); his effort was supported by the Coordinating Research Council, Inc. (CRC) through Contract No. A-129. The Coordinating Research Council, Inc. (CRC) is a non-profit corporation supported by the petroleum and automotive equipment industries. CRC operates through the committees made up of technical experts from industry and government who voluntarily participate. The four main areas of research within CRC are: air pollution (atmospheric and engineering studies); aviation fuels, lubricants, and equipment performance; heavy-duty vehicle fuels, lubricants, and equipment performance (e.g., diesel trucks); and light-duty vehicle fuels, lubricants, and equipment performance (e.g., passenger cars). CRC's function is to provide the mechanism for joint research conducted by the two industries that will help in determining the optimum combination of petroleum products and automotive equipment. CRC's work is limited to research that is mutually beneficial to the two industries involved. I.C. Faloona's effort was supported by the USDA National Institute of Food and Agriculture, (Hatch project CA-D-LAW-2481-H, "Understanding Background Atmospheric Composition, Regional Emissions, and Transport Patterns Across CA"). R.G. Derwent's efforts were pro bono.

Competing Interests

The contact authors have declared that none of the authors has any competing interests.

Author contributions

965 D.D.P. was responsible for the overall design. I.C.F. provided analysis. D.D.P. wrote the paper with input from I.C.F. and R.G.D. All authors: edited and revised manuscript.

Data availability statement

970 The California MDA8 ozone concentrations were obtained from the CA Air Resources Board archive (<https://www.arb.ca.gov/adam/index.html>; last accessed 24 October 2023). The ODVs were obtained from EPA's AQS data archive (<https://www.epa.gov/aqs>; last accessed 17 June 2022).

References

Abalos, M., Orbe, C., Kinnison, D. E., Plummer, D., Oman, L. D., Jöckel, P., Morgenstern, O., Garcia, R. R., Zeng, G., Stone, K. A., and Dameris, M.: Future trends in stratosphere-to-troposphere transport in CCMI models, *Atmos. Chem. Phys.*, 20, 6883–6901, <https://doi.org/10.5194/acp-20-6883-2020>, 2020.

Deleted: The ozone distribution in the troposphere results from the interactions of a great many chemical and physical processes driving ozone sources, sinks and transport, fully understanding this distribution requires that the impacts of each of those processes be accurately simulated and results fully integrated with each other. Chemical transport models (CTMs) are essential to this understanding; they approach this challenging task through computational parameterizations of each process in concert with the intention of reproducing observed characteristics of the ozone distribution. However, biases in present model results are quite often found to be in the range of 5-20 ppb (Hu et al., 2017; Zhang et al., 2020). Comparison of results between different model approaches demonstrates that large uncertainties remain in our understanding; Section S1 of the Supplement briefly discusses some model result comparisons. Such comparisons have led to the increasing recognition that CTMs are not yet able to provide accurate estimates of atmospheric ozone concentrations without imposing additional constraints directly from observations. For example, Skipper et al. (2021) present a method to fuse ozone observations with USB simulated by a regional CTM in order to correct model bias, and Hosseinpour et al. (2023) present a machine learning technique that improves CTM estimates of US background ODVs, which led to improved agreement with the Parrish et al. (2017) estimate for Los Angeles and the Parrish and Ennis (2019) estimate for New York City. The much simpler, observation-based model we use takes a contrasting approach; it avoids detailed process simulation by analyzing the measured ozone distribution, which necessarily reflects the accurately integrated impacts of all relevant processes, because the atmosphere itself has performed that integration. These results also have significant uncertainty, which arises from the underlying assumptions required to interpret observations so as to extract information regarding the processes that produced the observed ozone distribution. Notably, the simplicity of this model is helpful, because impacts of the model assumptions can be readily evaluated; Sections S2-S6 of the Supplement present some of these evaluations. In contrast, CTMs require a manifold of parameterizations to simulate the many relevant chemical and physical processes; these parameterizations are generally embedded within the model computer code, rendering it extremely difficult to evaluate the uncertainty that arises from any specific parameterization. For example, evaluations by Derwent et al. (2016; 2018; 2021) identify important uncertainties in CTM results, but determining avenues to correct the model deficiencies was not possible. Neither CTMs nor our simpler approach alone can give the desired full understanding of the tropospheric ozone distribution. ¶ To advance our understanding, we believe that two key points made by Held (2005) are particularly important: ¶ "It is fair to say that we typically gain some understanding of a complex system by relating its behavior to that of other, especially simpler, systems. For sufficiently complex systems, we need a model hierarchy on which to base our understanding, describing how the dynamics change as key sources of complexity are added or subtracted."¶ "Elegance versus Elaboration. An elegant model is only as elaborate as it needs to be to capture the essence of a particular source of complexity, but is no more elaborate."¶ Emanuel (2020) makes similar points from a somewhat different perspective: ¶ "As it becomes easier to undertake complex computer simulations of climate and weather, ... it is tempting to use computers to simulate, rather than understand, nature." ¶ (... [1])

- Abeleira, A., and Farmer, D.: Summer ozone in the Northern Front Range Metropolitan Area: Weekend–weekday effects, temperature dependences, and the impact of drought. *Atmospheric Chemistry and Physics*, 17: 6517–6529. DOI: <http://dx.doi.org/10.5194/acp-17-6517-2017>, 2017.
- 1110 Ahmadi, M., and John, K.: Statistical evaluation of the impact of shale gas activities on ozone pollution in North Texas. *Science of The Total Environment*, 536, 457–467, 2015.
- Berlin, S.R., Langford, A.O., Estes, M., Dong, M., and Parrish, D.D.: Magnitude, decadal changes and impact of regional background ozone transported into the greater Houston, Texas area, *Environ. Sci. Technol.*, 47(24), 13985–13992, doi:10.1021/es4037644, 2013.
- 1115 Burke, M., Driscoll, A., Heft-Neal, S., Xue, J., Burney, J. and Wara, M.: The changing risk and burden of wildfire in the United States, *Proceedings of the National Academy of Sciences*, doi:10.1073/pnas.2011048118, 2021.
- Cooper, O.R., Oltmans, S.J., Johnson, B.J., Brioude, J., Angevine, W., Trainer, M., Parrish, D.D., Ryerson, T.R., Pollack, I., Cullis, P.D., Ives, M.A., Tarasick, D.W., Al-Saadi, J. and Stajner, I.: Measurement of western U.S. baseline ozone from the surface to the tropopause and assessment of downwind impact regions, *J. Geophys. Res.*, 116, D00V03, doi:10.1029/2011JD016095, 2011.
- 1120 Cooper, O. R., Parrish, D. D., Ziemke, J., Balashov, N. V., Cupeiro, M., Galbally, I. E., Gilge, S., Horowitz, L., Jensen, N. R., Lamarque, J.-F., Naik, V., Oltmans, S. J., Schwab, J., Shindell, D. T., Thompson, A. M., Thouret, V., Wang, Y., and Zbinden, R. M.: Global distribution and trends of tropospheric ozone: An observation-based review, *Elem. Sci. Anth.* 2: 000029 doi: 10.12952/journal.elementa.000029, 2014.
- 1125 Cooper, O. R., Langford, A. O., Parrish, D. D., and Fahey, D. W.: Challenges of a lowered US ozone standard, *Science*, 348(6239), 1096–1097, doi:10.1126/science.aaa5748, 2015.
- Cooper, O. R., Schultz, M. G., Schröder, S., Chang, K.-L., Gaudel, A., Benítez, G. C., Cuevas, E., Fröhlich, M., Galbally, I. E., Molloy, S., Kubistin, D., Lu, X., McClure-Begley, A., Nédélec, P., O'Brien, J., Oltmans, S. J., Petropavlovskikh, I., 1130 Ries, L., Senik, I., Sjöberg, K., Solberg, S., Spain, G. T., Spangl, W., Steinbacher, M., Tarasick, D., Thouret, V., and Xu, X.: Multi-decadal surface ozone trends at globally distributed remote locations. *Elem. Sci. Anth.*, 8(23), DOI: <https://doi.org/10.1525/elementa.420>, 2020.
- Derwent, R. G., Parrish, D. D., Galbally, I. E., Stevenson, D. S., Doherty, R. M., Young, P. J., and Shallcross, D. E.: Interhemispheric differences in seasonal cycles of tropospheric ozone in the marine boundary layer: Observation-model comparisons, *J. Geophys. Res. Atmos.*, 121, 11,075–11,085, doi:10.1002/2016JD024836, 2016.
- 1135 Derwent, R. G., Parrish, D. D., Galbally, I. E., Stevenson, D. S., Doherty, R. M., Naik, V., and Young, P. J.: Uncertainties in models of tropospheric ozone based on Monte Carlo analysis: Tropospheric ozone burdens, atmospheric lifetimes and surface distributions, *Atmos. Environ.*, 180, 93–102, doi.org/10.1016/j.atmosenv.2018.02.047, 2018.
- Derwent, R. G., Parrish, D. D., Archibald, A. T., Deushi, M., Bauer, S. E., Tsigaridis, K., Shindell, D., Horowitz, L. W., 1140 Anwar, M., Khan, H., and Shallcross D. E.: Intercomparison of the representations of the atmospheric chemistry of pre-

- industrial methane and ozone in earth system and other global chemistry-transport models, *Atmos. Environ.*, 248, <https://doi.org/10.1016/j.atmosenv.2021.118248>, 2021.
- Derwent, R. G., Parrish, D. D., and Faloon, I. C.: Opinion: Establishing a science-into-policy process for tropospheric ozone assessment, *Atmos. Chem. Phys.*, 23, 13613–13623, <https://doi.org/10.5194/acp-23-13613-2023>, 2023.
- 1145 Dolwick, P., Akhtar, F., Baker, K. R., Possiel, N., Simon, H., and Tonnesen, G.: Comparison of background ozone estimates over the western United States based on two separate model methodologies, *Atmos. Environ.*, 109, 282–296, <https://doi.org/10.1016/j.atmosenv.2015.01.005>, 2015.
- Emanuel, K.: The Relevance of Theory for Contemporary Research in Atmospheres, Oceans, and Climate. *AGU Advances*, 1, e2019AV000129, doi: <https://doi.org/10.1029/2019AV000129>, 2020.
- 1150 Evans, J. M., and Helmig, D.: Investigation of the influence of transport from oil and natural gas regions on elevated ozone levels in the northern Colorado front range, *Journal of the Air & Waste Management Association*, 67:2, 196-211, DOI:10.1080/10962247.2016.1226989, 2017.
- Faloon, I. C., Chiao, S., Eiserloh, A. J., Alvarez II, R. J., Kirgis, G., Langford, A. O., Senff, C. J., Caputi, D., Hu, A., Iraci, L. T., Yates E. L., Marrero, J. E., Ryoo, J.-M., Conley, S., Tanrikulu, S., Xu, J., and Kuwayama, T.: The California baseline ozone transport study (CABOTS). *Bull. Amer. Meteor. Soc* 101 (4):E427–E445. doi:10.1175/BAMS-D-18-0302.1, 2020.
- 1155 Fiore, A. M., Oberman, J. T., Lin, M. Y., Zhang, L., Clifton, O. E., Jacob, D. J., Naik, V., Horowitz, L. W., Pinto, J. P., and Milly, G. P.: Estimating North American background ozone in U.S. surface air with two independent global models: Variability, uncertainties, and recommendations, *Atmos. Environ.*, 96, 284-300, doi.org/10.1016/j.atmosenv.2014.07.045, 2014.
- 1160 Flynn, M. T., Mattson, E. J., Jaffe, D.A., and Gratz, L. E.: Spatial patterns in summertime surface ozone in the Southern Front Range of the U.S. Rocky Mountains. *Elem Sci Anth*, 9: 1. DOI: <https://doi.org/10.1525/elementa.2020.00104>, 2021.
- Geddes, J. A., Pusede, S. E., and Wong, A. Y. H.: Changes in the relative importance of biogenic isoprene and soil NO_x emissions on ozone concentrations in nonattainment areas of the United States, *Journal of Geophysical Research: Atmospheres*, 127, e2021JD036361. <https://doi.org/10.1029/2021JD036361>, 2022.
- 1165 Gkatzelis, G. I., Gilman, J. B., Brown, S. S., Eskes, H., Gomes, A. R., Lange, A. C., and Kiendler-Scharr, A.: The global impacts of COVID-19 lockdowns on urban air pollution: A critical review and recommendations. *Elementa Science of the Anthropocene*, 9(1). <https://doi.org/10.1525/elementa.2021.00176>, 2021.
- 1170 Gong, X., Kaulfus, A., Nair, U. and Jaffe, D. A.: Quantifying O₃ impacts in urban areas due to wildfires using a generalized additive model, *Environ. Sci. Technol.*, 51, 22, 13216–13223, <https://doi.org/10.1021/acs.est.7b03130>, 2017.
- Held, I. M.: The gap between simulation and understanding in climate modeling. *Bull. Am. Meteorol. Soc.*, 86, 1609-1614, doi:10.1175/Bams-86-11-1609, 2005.

- 1175 Hogrefe C., Liu, P., Pouliot, G., Mathur, R., Roselle, S., Flemming, J., Lin, Me., and Park, R. J.: Impacts of different characterizations of large-scale background on simulated regional-scale ozone over the continental United States, *Atmos. Chem. Phys.*, 18, 3839–3864, <https://doi.org/10.5194/acp-18-3839-2018>, 2018.
- Hosseinpour, F., Kumar, N., Tran, T., and Knipping, E.: Using machine learning to improve the estimate of U.S. background ozone, *Atmospheric Environment*. 316. 120145. 10.1016/j.atmosenv.2023.120145, 2023.
- 1180 Hu, L., Jacob, D. J., Liu, X., Zhang, Y., Zhang, L., Kim, P. S., Sulprizio, M. P., and Yantosca R. M.: Global budget of tropospheric ozone: Evaluating recent model advances with satellite (OMI), aircraft (IAGOS), and ozonesonde observations, *Atmos. Environ.*, 167, 323-334, 2017.
- Iglesias, V., Balch, J. K., and Travis, W. R.: U.S. fires became larger, more frequent, and more widespread in the 2000s, *Sci. Adv.* 8, eabc0020, 2022.
- 1185 Jaffe, D., Price, H., Parrish, D. D., Goldstein, A., and Harris, J.: Increasing background ozone during spring on the west coast of North America, *Geophysical Research Letters*, 30 (12), 1613, doi:10.1029/2003GL017024, 2003.
- Jaffe, D. A., and Ray, J.: Increase in Ozone at Rural Sites in the Western U.S., *Atmos. Environ.*, 41, 5452–5463, 2007.
- Jaffe, D. A., Cooper, O. R., Fiore, A. M., Henderson, B. H., Tonneson, G. S., Russell, A. G., Henze, D. K., Langford, A. O., Lin, M., and Moore, T.: Scientific assessment of background ozone over the U.S.: Implications for air quality management, *Elem. Sci. Anth.*, 6, 56, <https://doi.org/10.1525/elementa.309>, 2018.
- 1190 Karle, N. N., Fitzgerald, R. M., Sakai, R. K., Sullivan, D. W., and Stockwell, W. R.: Multi-Scale Atmospheric Emissions, Circulation and Meteorological Drivers of Ozone Episodes in El Paso-Juarez Airshed, *Atmosphere*, 12(12), doi:ARTN 1575 10.3390/atmos12121575, 2021.
- Kim, S.-W., McDonald, B. C., Seo, S., Kim, K.-M., and Trainer, M.: Understanding the paths of surface ozone abatement in the Los Angeles Basin. *Journal of Geophysical Research: Atmospheres*, 127, e2021JD035606. <https://doi.org/10.1029/2021JD035606>, 2022.
- 1195 Langford, A. O., Alvarez II, R. J., Brioude, J., Fine, R., Gustin, M. S., Lin, M. Y., Marchbanks, R. D., Pierce, R. B., Sandberg, S. P., Senff, C. J., Weickmann, A. M., and Williams, E. J.: Entrainment of stratospheric air and Asian pollution by the convective boundary layer in the southwestern U.S., *J. Geophys. Res.*, 122(2), 1312-1337, doi:10.1002/2016JD025987, 2017.
- 1200 Langford, A. O., Senff, C. J., Alvarez II, R. J., Aikin, K. C., Baidar, S., Bonin, T. A., Brewer, A. W., Brioude, J., Brown, S. S., Burley, J. D., Caputi, D. J., Conley, S. A., Cullis, P. D., Decker, Z., Evan, S., Kirgis, G., Lin, M., Pagowski, M., Peischl, J., Petropavlovskikh, I., Pierce, R. B., Ryerson, T. B., Sandberg, S. P., Sterling, C. W., Weickmann, A. M., and Zhang, L.: The Fires, Asian, and Stratospheric Transport–Las Vegas Ozone Study (FAST-LVOS), *Atmos. Chem. Phys.*, 22(3), 1707-1737, doi:10.5194/acp-22-1707-2022.ee, 2022.
- 1205 Lin, M., Fiore, A. M., Horowitz, L. W., Cooper, O. R., Naik, V., Holloway, J., Johnson, B. J., Middlebrook, A. M., Oltmans, S. J., Pollack, I. B., Ryerson, T. B., Warner, J. X., Wiedinmyer, C., Wilson, J., and Wyman, B.: Transport of Asian ozone

- pollution into surface air over the western United States in spring, *J. Geophys. Res.*, 117, D00V07, <https://doi.org/10.1029/2011jd016961>, 2012a.
- 1210 Lin, M., Fiore, A. M., Cooper, O. R., Horowitz, L. W., Langford, A. O., Levy, H., Johnson, B. J., Naik, V., Oltmans, S. J., and Senff, C. J.: Springtime high surface ozone events over the western United States: Quantifying the role of stratospheric intrusions, *J. Geophys. Res.*, 117, D00V22, <https://doi.org/10.1029/2012jd018151>, 2012b.
- Lin, M., Horowitz, L. W., Payton, R., Fiore, A. M., and Tonnesen, G.: US surface ozone trends and extremes from 1980 to 2014: quantifying the roles of rising Asian emissions, domestic controls, wildfires, and climate, *Atmos. Chem. Phys.*, 17, 2943–2970, <https://doi.org/10.5194/acp-17-2943-2017>, 2017.
- 1215 Lu, Z., Streets, D. G., de Foy, B., Lamsal, L. N., Duncan, B. N., and Xing, J.: Emissions of nitrogen oxides from US urban areas: estimation from Ozone Monitoring Instrument retrievals for 2005–2014, *Atmos. Chem. Phys.*, 15, 10367–10383, www.atmos-chem-phys.net/15/10367/2015/ doi:10.5194/acp-15-10367-2015, 2015.
- McDonald, B. C., de Gouw, J. A., Gilman, J. B., Jathar, S. H., Akherati, A., Cappa, C. D., Jimenez, J. L., Lee-Taylor, J., Hayes, P. L., McKeen, S. A., Cui, Y. Y., Kim, S.-W., Gentner, D. R., Isaacman-VanWertz, G., Goldstein, A. H., Harley, 1220 R. A., Frost, G. J., Roberts, J. M., Ryerson, T. B., and Trainer, M.: Volatile chemical products emerging as largest petrochemical source of urban organic emissions, *Science*, 359, 760–764, 2018.
- McDuffie, E. E., Edwards, P. M., Gilman, J. B., Lerner, B. M., Dubé, W. P., Trainer, M., Wolfe, D. E., Angevine, W. M., deGouw, J., Williams, E. J., Tevlin, A. G., Murphy, J. G., Fischer, E. V., McKeen, S., Ryerson, T. B., Peischl, J., Holloway, J. S., Aikin, K., Langford, A. O., Senff, C. J., Alvarez, R. J., Hall, S. R., Ullmann, K., Lantz, K. O., Brown, S.: 1225 S.: Influence of oil and gas emissions on summertime ozone in the Colorado Northern Front Range, *J. Geophys. Res.: Atmos.*, 121, 8712–8729, 2016.
- McKeen, S. A., Wotawa, G., Parrish, D. D., Holloway, J. S., Burh, M. P., Hübler, G., Fehsenfeld, F. C., and Meagher, J. F.: Ozone production from Canadian wildfires during June and July of 1995, *J. Geophys. Res.*, 107 (D14):4192. doi:10.1029/2001JD000697, 2002.
- 1230 Nopmongcol, U., Alvarez, Y., Jung, J., Grant, J., Kumar, N., and Yarwood, G.: Source contributions to United States ozone and particulate matter over five decades from 1970 to 2020, *Atmos. Environ.*, 167, 116–128, 2017.
- Oltmans, S. J., Lefohn, A. S., Harris, J. M., and Shadwick, D. S.: Background ozone levels of air entering the west coast of the U.S. and assessment of longer-term changes. *Atmos. Environ* 42 (24):6020–6038, doi:10.1016/j.atmosenv.2008.03.034, 2008.
- 1235 Parrish, D. D., Millet, D. B., and Goldstein, A.H.: Increasing ozone in marine boundary layer air inflow at the west coasts of North America and Europe, *Atmospheric Chemistry and Physics*, 9, 1303–1323, 2009.
- Parrish, D. D., Aikin, K. C., Oltmans, B. J., Johnson, S. J., Ives, M., and Sweeny, C. (2010), Impact of transported background ozone inflow on summertime air quality in a California ozone exceedance area, *Atmos. Chem. Phys.*, 10, 10093–10109, doi:10.5194/acp-10-10093-2010.

- 1240 Parrish, D. D., Law, K. S., Staehelin, J., Derwent, R., Cooper, O. R., Tanimoto, H., Volz-Thomas, A., Gilge, S., Scheel, H.-E., Steinbacher, M., and Chan, E.: Long-term changes in lower tropospheric baseline ozone concentrations at northern mid-latitudes, *Atmos. Chem. Phys.*, 12, 11,485–11,504, 2012.
- Parrish, D. D. and Stockwell, W. R.: Urbanization and air pollution: Then and now, *Eos: Earth & Space Science News*, 96, 10–15, 2015.
- 1245 Parrish, D.D., Xu, J., Croes, B. and Shao, M.: Air Quality Improvement in Los Angeles - Perspectives for Developing Cities, *Frontiers of Environ Sci. & Engineering*, 10(5): 11, doi: 10.1007/s11783-016-0859-5, 2016.
- Parrish, D. D., Young, L. M., Newman, M. H., Aikin, K. C., and Ryerson, T. B.: Ozone design values in southern California's air basins: Temporal evolution and U.S. background contribution, *J. Geophys. Res.*, 122, 11,166–11,182, 2017.
- 1250 Parrish, D. D., and Ennis, C.A.: Estimating background contributions and US anthropogenic enhancements to maximum ozone concentrations in the northern US, *Atmos. Chem. Phys.*, 19, 12587–12605, <https://doi.org/10.5194/acp-19-12587-2019>, 2019.
- Parrish, D. D., Derwent, R. G., O'Doherty, S. and Simmonds, P. G.: Flexible approach for quantifying average long-term changes and seasonal cycles of tropospheric trace species, *Atmos. Meas. Tech.*, 12, 3383–3394, <https://doi.org/10.5194/amt-12-3383-2019>, 2019.
- 1255 Parrish, D. D., Derwent, R. G., Steinbrecht, W., Stübi, R., VanMalderen, R., Steinbacher, M., Trickl, T., Ries, L., and Xu, X.: Zonal similarity of long-term changes and seasonal cycles of baseline ozone at northern midlatitudes. *J. Geophys. Res.: Atmos.*, doi: 10.1029/2019JD031908, 2020.
- Parrish, D.D., Derwent, R.G., and Faloon, I.C.: Long-term baseline ozone changes in the Western US: A synthesis of analyses, *Journal of the Air & Waste Management Association*, DOI: 10.1080/10962247.2021.1945706, 2021.
- 1260 Parrish, D.D., Derwent, R.G., and Faloon, I.C.: Observational-based assessment of contributions to maximum ozone concentrations in the western US, *Journal of the Air & Waste Management Association*, 72:5, 434–454, <https://doi.org/10.1080/10962247.2022.2050962>, 2022.
- Schnell, R. C., Oltmans, S. J., Neely, R. R., Endres, M. S., Molenaar, J. V., and White, A. B.: Rapid photochemical production of ozone at high concentrations in a rural site during winter, *Nature Geosci.* 2, 120–122, 2009.
- 1265 Simon, H., Reff, A., Wells, B., Xing, J., and Frank, N.: Ozone trends across the United States over a period of decreasing NO_x and VOC emissions. *Environmental Science & Technology*, 49(1), 186–195. <https://doi.org/10.1021/es504514z>, 2015.
- Skerlak, B., Sprenger, M., and Wernli, H.: A global climatology of stratosphere-troposphere exchange using the ERA-Interim data set from 1979 to 2011, *Atmos. Chem. Phys.* 14, 913–937, 2014.
- 1270 Skipper, T. N., Hu, Y., Odman, M. T., Henderson, B. H., Hogrefe, C., Mathur, R., Russell, A. G.: Estimating US Background Ozone Using Data Fusion, *Environmental Science and Technology*, 55, 4504–4512, DOI: 10.1021/acs.est.0c08625, 2021.

- Sprenger, M., and Wernli, H.: A northern hemisphere climatology of cross-tropopause exchange for the ERA15 time period (1979-1993), *J. Geophys. Res.*, 108, 8521-, <https://doi.org/10.1029/2002JD002636>, 2003.
- 1275 Westerling, A. L., Hidalgo, H. G., Cayan, D. R., and Swetnam, T. W.: Warming and earlier spring increase western US forest wildfire activity, *Science*, 313, 940-943, 2006.
- Westerling, A. L.: Increasing western US forest wildfire activity: sensitivity to changes in the timing of spring, *Philos Trans R Soc Lond B Biol Sci.* 2016 Jun 5;371(1696):20150178. doi: 10.1098/rstb.2015.0178, 2016.
- Yates, E. L., Iraci, L. T., Austerberry, D., Pierce, R. B., Roby, M. C., Tadić, J. M., Loewenstein, M., and Gore W.:
1280 Characterizing the impacts of vertical transport and photochemical ozone production on an exceedance area, *Atmos. Environ.*, 109, 342-350, 2015.
- Zhang, J., Yongjie, W., and Zhangfu, F.: Ozone Pollution: A Major Health Hazard Worldwide, *Front. Immunol.*, 10, <https://doi.org/10.3389/fimmu.2019.02518>, 2019.
- Zhang, L., Lin, M., Langford, A. O., Horowitz, L. W., Senff, C. J., Klovenski, E., Wang, Y., Alvarez II, R. J., Petropavlovskikh,
1285 I., Cullis, P., Sterling, C. W., Peischl, J., Ryerson, T. B., Brown, S. S., Decker, Z. C. J., Kirgis, G., and Conley, S.:
Characterizing sources of high surface ozone events in the southwestern US with intensive field measurements and two
global models, *Atmos. Chem. Phys.*, 20, 10379–10400, <https://doi.org/10.5194/acp-20-10379-2020>, 2020.
- Ziemke, J. R., Chandra, S., Labow, G. J., Bhartia, P. K., Froidevaux, L., and Witte, J. C.: A global climatology of tropospheric
and stratospheric ozone derived from Aura OMI and MLS measurements, *Atmos. Chem. Phys.*, 11, 9237–9251,
1290 <https://doi.org/10.5194/acp-11-9237-2011>, 2011.

Table 1. Parameter values (with 95% confidence limits) derived from fits of Equation 1 to time series baseline ozone concentrations. The d parameter represents the coefficient of the cubic term in the analogous fits to a third order polynomial.

Data Set	b (ppb yr ⁻¹)	c (10 ⁻² ppb yr ⁻²)	year _{max}	RMSD (ppb)	Reference	d (10 ⁻⁴ ppb yr ⁻³)
2-year means of baseline ozone	0.20 ± 0.06	-1.8 ± 0.6	2005.7 ± 2.5	1.4	Parrish et al., 2020	+0.6 ± 5.7
CASTNET ODVs – normalized ¹	0.07 ± 0.13	-1.5 ± 0.8	2002.1 ± 4.4	2.4	This work	+5.8 ± 10.1

¹ The a parameter value derived from the fit of Equation 1 to the normalized CASTNET ODVs is 71.3 ± 0.8 ppb, which is consistent with the normalization process utilized here; the corresponding value in the Parrish et al. (2020) analysis was near zero due to the different normalization approach used in that work.

1295

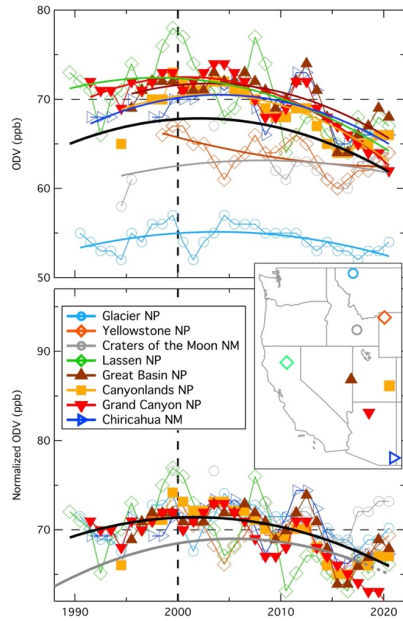


Figure 1: Time series of ODVs recorded at the eight rural western CASTNET sites shown in the inset map. Solid symbols indicate the three sites used for the normalization. Solid curves indicate the fits of Equation 1 to the individual site data (upper panel, including the black curve fit to all data) and to all normalized data (black curve, lower panel). The lower grey curve in the lower panel is the Parrish et al. (2020) fit to baseline data, normalized to 68.5 ppb (the a parameter value derived in a fit of Equation 3 to the normalized CASTNET data) in the year 2000; dashed line shows extrapolation of that fit to 2021.

1300

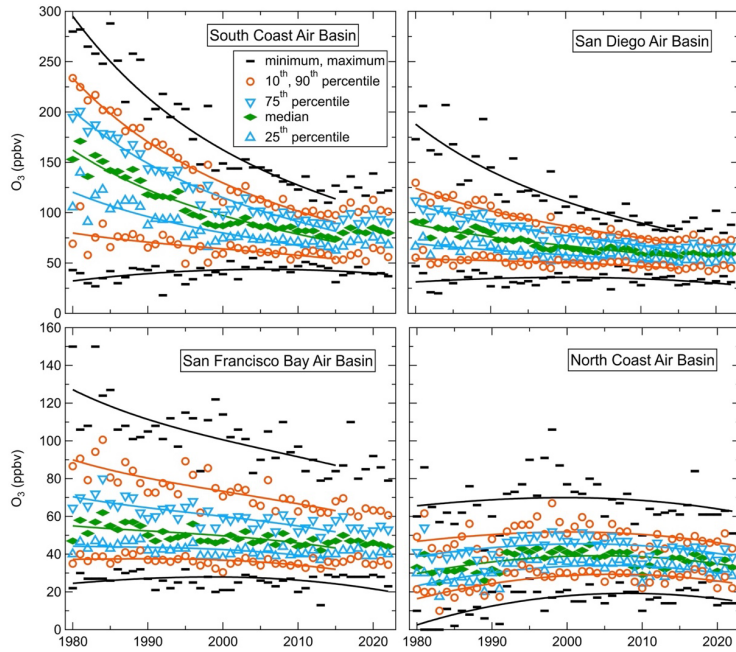
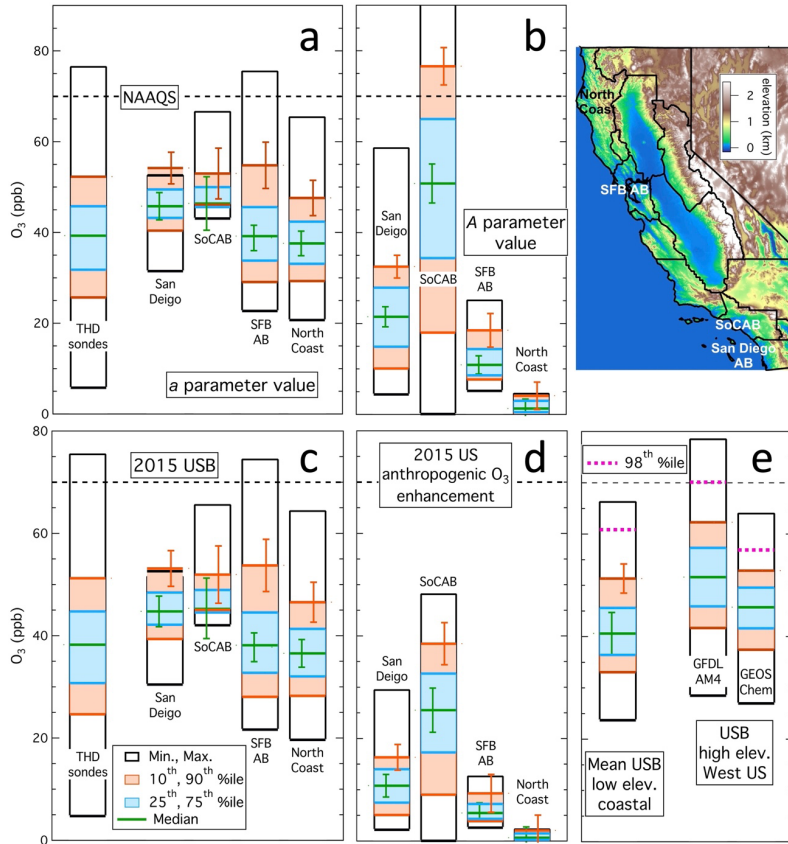
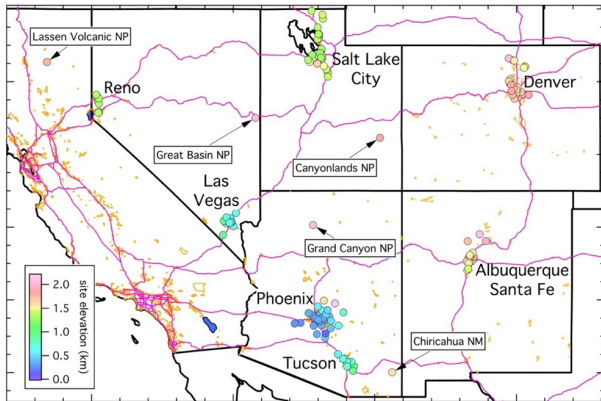


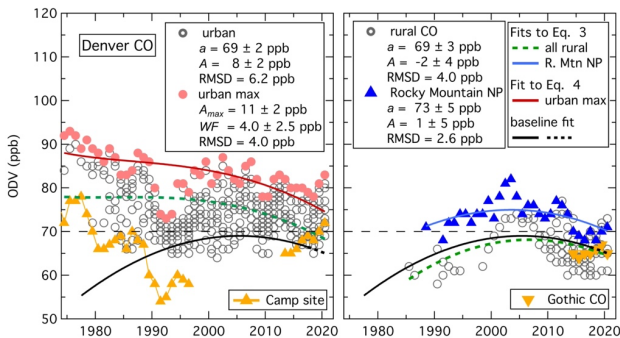
Figure 2: Temporal evolution of distribution of MDA8 ozone concentrations in four CA air basins. Symbols indicate the annotated percentiles of the annual distribution of the largest MDA8 concentration recorded at any monitor within the basin on each day of the May–September ozone season. Solid curves indicate fits of Equation 3 to the respective percentiles.



1310 **Figure 3.** Comparisons of estimates of MDA8 ozone component distributions in 4 coastal California air basins; the inset
 1315 map identifies the 4 air basins, with the black lines indicating air basin boundaries. **(a)** Comparison of ozone concentrations
 measured between 0.6 and 1 km altitude by sondes launched from Trinidad Head from May through September, 1997-2017
 and derived *a* parameter values. **(b)** Comparison of derived *A* parameter values. **(c)** Comparison of the same sonde ozone
 measurements and year 2015 USB distribution. **(d)** Comparison of year 2015 distributions of US anthropogenic
 enhancements. **(e)** Comparison of the mean of the five, year 2015 USB distributions illustrated in (c) with the distributions of
 daily MDA8 USB for April–June 2017 at western US high-elevation sites simulated by two global models; these latter bar
 graphs were derived from Figure 17 of Zhang et al. (2020). Error bars on the median and 90th percentile lines indicate
 estimated uncertainties of the determination of the USB and the US anthropogenic ozone enhancement percentiles, and the
 standard deviations of the means from the five USB determinations.

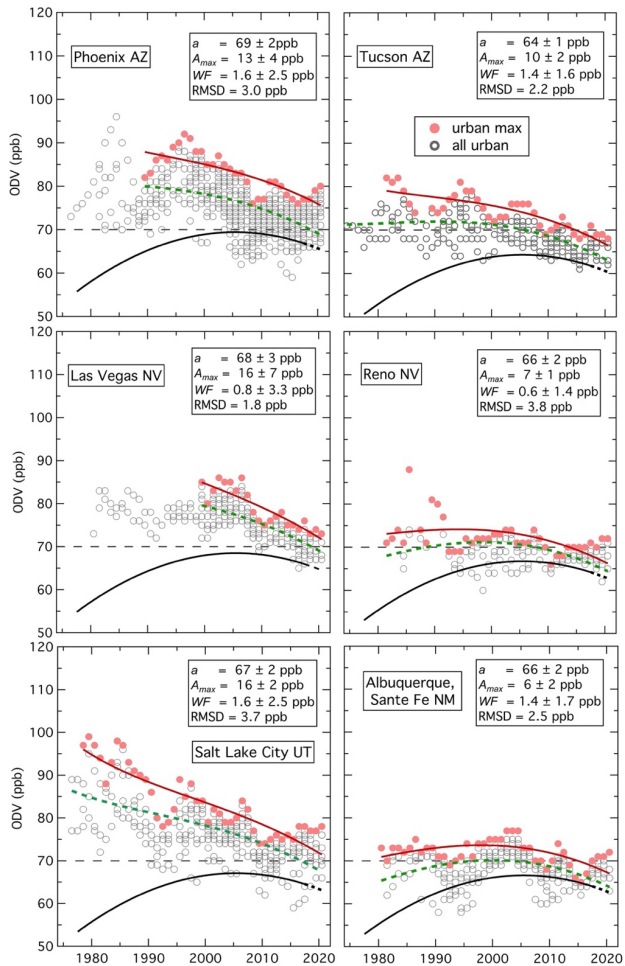


1320 **Figure 4:** Map of southwestern US urban monitoring sites; the symbols are color-coded according to site elevation as indicated in the annotation. Lines indicate outlines of southwestern US states (black), urban areas (gold) and interstates and selected other major highways (violet). Seven urban areas, whose sites are analyzed together as separate data sets, are labelled. Locations of five of the isolated rural CASTNET sites are also included.

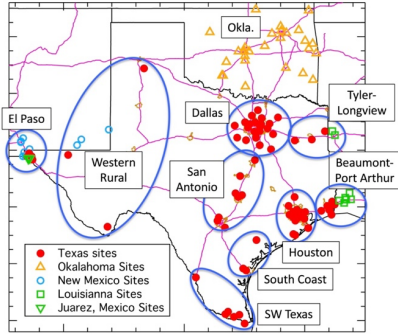


1325 **Figure 5:** Time series of ODVs recorded in Colorado at the urban sites (left graph) whose locations are shown in Figure 4, and at rural sites (right graph, excluding the Four Corners area) whose locations are shown in Figure S1. Open symbols indicate all ODVs recorded in each area, and solid symbols indicate ODVs from three specific sites and the maximum Denver urban ODVs. As indicated by an annotation on the right graph, green dashed curves indicate fits of Equation 3 to all plotted ODVs (excluding the Camp and Rocky Mountain NP sites, blue curve on right indicates Rocky Mountain NP fit of Equation 3, and red solid curve on left indicates fit of Equation 4 to the maximum Denver urban ODVs; parameters derived in these fits are annotated in the graphs. The black solid curves with dashed extensions indicate the fit to the baseline data included as the grey curve in Figure 1. The light dashed lines indicate the 70 ppb ozone NAAQS.

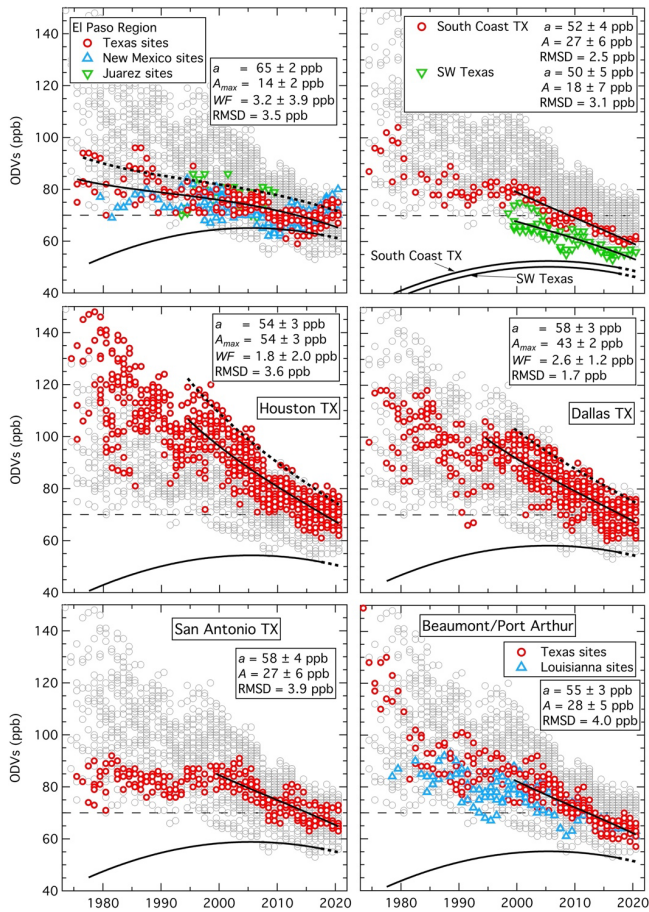
1330



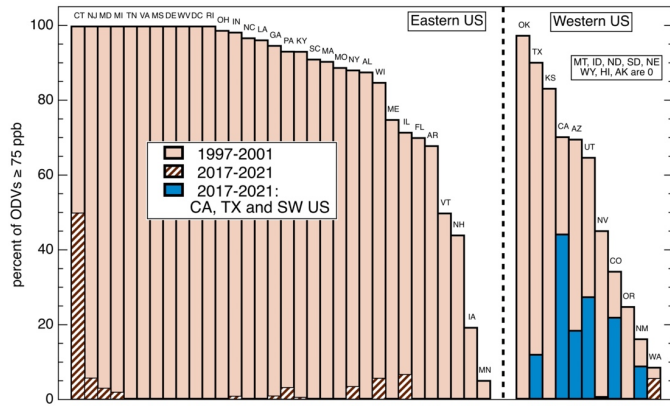
1335 **Figure 6:** Time series of ODVs recorded in six southwestern US urban areas shown in Figure 4. Open symbols indicate all ODVs recorded in each area, and the solid symbols indicate the maximum ODVs in each year in each area. As in Figure 5 green dashed curves indicate fits of Equation 3 to all ODVs; red solid curves indicate fits of Equation 4 to the maximum ODVs, with the parameters derived in this fit annotated. The black solid curves with dashed extensions indicate the fit to the baseline data from Figure 1, normalized to the respective a parameter values. The light dashed lines indicate the 70 ppb ozone NAAQS.



1340 **Figure 7:** Map of monitoring sites within the Texas region; the symbols are color-coded according to state as indicated in the annotation. Lines indicate outlines of southwestern US states (black), urban areas (gold) and interstates (violet). Nine Texas regions, whose sites are analyzed together as separate data sets, are indicated.



1345 **Figure 8:** Time series of ODVs recorded in seven of the Texas regions shown in Figure 7. Grey symbols in each graph indicate all recorded Texas ODVs. Colored symbols indicate the ODVs from each respective region. Upper black solid curves indicate fits of Equation 3 to all ODVs in the area over the curve's time span; the parameters derived in these fits are annotated in three graphs; dotted curves indicate fits of Equation 4 to the maximum ODVs in El Paso, Houston and Dallas, with the parameters derived in these fits annotated. Lower solid curves with dashed extensions indicate the fit to the baseline data from Figure 1, normalized to the respective a parameter values. The light dashed lines indicate the 70 ppb ozone NAAQS.



1350

Figure 9: Comparison of percentage of ODVs greater than or equal to 75 ppb recorded at all sites in individual states over two 5-year periods: 2017-2021 (hatched and dark blue bars) and a period 20 years earlier - 1997-2001 (light-colored bars). Individual states are indicated by their two letter abbreviations (defined in Table S6). States are arbitrarily divided between eastern and western regions. Southwestern states, Texas and California are indicated by solid dark blue bars. Eight rural states, all in the western region, reported no ODVs greater than or equal to 75 ppb in either period.

1355

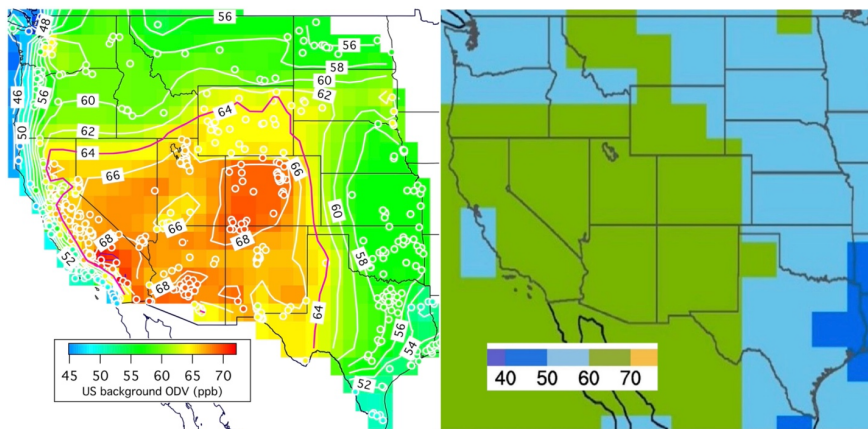
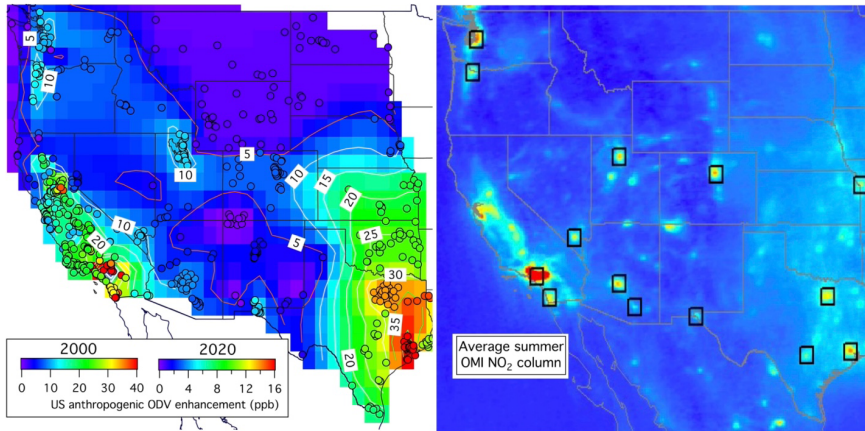


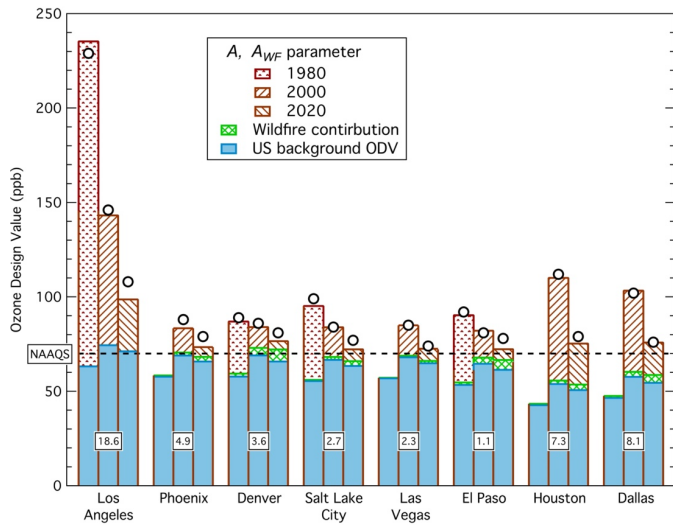
Figure 10: Two estimates of the spatial variability of US background ODVs over the western US. **(Left)** Contour map of estimated US background ODV for year 2000. The US background ODV excludes estimated wildfire contributions. The 64 ppb contour is colored violet to indicate the extensive area of largest values. Symbols in the contour map indicate individual monitoring sites included in the analysis with the same color coding as the contour map. Results from Parrish et al. (2017; 2022) and estimates for California’s Central Valley (Faloona et al., manuscript in preparation) are included. **(Right)** Annual 4th highest MDA8 ozone concentration averaged over 2010–2014, from a GFDL-AM3 model simulation with North American anthropogenic emissions zeroed out (figure reproduced from a section of Figure 3 of Jaffe et al., 2018).

1360

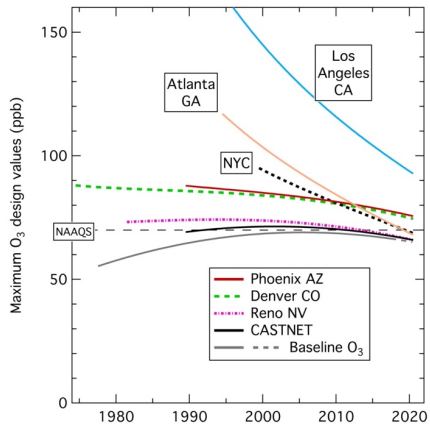
1365



1370 **Figure 11:** US anthropogenic ODV enhancement over the western US compared to mean summertime OMI NO₂ columns.
 1375 **(Left)** Contour map of estimated US anthropogenic ODV enhancement for year 2000 over the western US. Symbols in the contour map indicate individual monitoring sites included in the analysis with the same color coding as the contour map. A second color scale is included for interpretation of the colors as the year 2020 US anthropogenic ODV enhancement. Results from Parrish et al. (2017; 2022) and estimates for California's Central Valley (Faloona et al., manuscript in preparation) are included. **(Right)** The OMI NO₂ columns measured in April – September 2005-2014 are reproduced from Figure 1a of Lu et al. (2015) by cropping and changing the aspect ratio of that figure to approximate that of the contour map.



1380 **Figure 12:** Derived apportionment of maximum ODVs for 3 years at 20-year intervals for the Crestline site in Los Angeles,
 1385 four southwestern US and three Texas urban areas. The lower, solid blue bars indicate the US background ODV; they exclude
 estimated wildfire contributions, which are separately indicated by the middle, cross-hatched green bar segments. The top,
 brown patterned bars indicate the estimated US anthropogenic ODV enhancements, again excluding the wildfire contribution.
 For four cities the US anthropogenic ODV enhancements and wildfire contributions are missing in 1980 since the tabulated
 ODVs are inadequate to estimate that parameter that early in the monitoring record. The 2020 populations of the urban areas
 are annotated in millions. The circles indicate the actual maximum ODVs recorded in the respective years in each area.



1390 **Figure 13:** Comparison of fits of Equation 4 to time series of maximum ODVs recorded in three of the nation’s largest urban areas and three southwestern urban areas, with the fits to normalized rural western CASTNET ODVs and baseline ozone from Figure 1. The fitted urban curves are taken from Figure 2 of Parrish et al. (2022) for Los Angeles, Figure S6 for New York City (i.e., NYC) and Atlanta GA, Figure 5 for Denver CO, Figure 6 for Phoenix AZ and Reno NV, and Figure 1 for CASTNET and baseline ozone.