## Technical Corrections for EGUSPHERE-2024-1460

## May 12, 2025

## **Technical Corrections**

We thank the editor for their work overseeing the review of this manuscript and for their suggested technical corrections for improving the accepted manuscript prior to publication. We have implemented them all as detailed below. We also attach a differenced version of the manuscript from the accepted version to fully document the requested corrections.

- 1. Is there any specific meaning to the grid that is superimposed on Figure 1? If they coincide with lat/lon coordinates, none are indicated. Nor does the grid size line up with the "10 km" scale bar.
  - We have added the lat lon labels to Figure 1.
- 2. Clarify "nearest neighbors" in Line 119. Do you mean only the overlying pixel and its single nearest neighbor? Or more than a single nearest neighbor?
  - We have clarified the text to read "the TROPOMI pixel over Whittier and the eight directly adjacent pixels (nearest neighbors) were averaged"
- 3. You mention fitting uncertainties in the MAX-DOAS retrieval. Are these the only uncertainties that lead to the uncertainty bars in Figure 3 and 4? Or are other components of uncertainty included in these? What uncertainty product is used for the TROPOMI uncertainty bars?
  - In line 122 of the accepted manuscript we defined the errors used as follows "Error bars for the MAX-DOAS retrievals reflect propagated dSCD measurement errors and a priori error calculated using HEIPRO. Error bars for the TROPOMI observations reflect the spatial variability over the nearest neighbor pixels as calculated by the standard error of the mean." In response to the editor's suggestion we have added this information to the captions of Fig.3 and 4 as well.
- 4. Consider rewording: "NO2 vertical distributions tend to be more surface based events..." in Line 145 (it's awkward and a bit unclear)
  - We have revised surface based events to read "have more of the column in the lowest 200 m of the atmosphere".

# Measurement Report: Diurnal Variability of NO<sub>2</sub> and HCHO Lower Tropospheric Vertical Profiles in Southeastern Los Angeles

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Abstract. Ground level ozone in excess of United States ambient air quality standards remains a prevalent issue across Southern California, particularly in the summer months. To improve our understanding of the vertical distribution of ozone precursors in southern California, we used ground-based MAX-DOAS measurements in Whittier CA to simultaneously retrieve both near-surface mole fractions and vertical column densities of both NO2 and HCHO. Ratios of HCHO to NO2, commonly referred to as FNR, derived from satellite-based measurements are used to diagnose ozone production chemistry over regions without consistent surface based measurements. While vertical column densities of NO2 are well correlated with TROPOMI observations over the study period (R=0.73), HCHO VCDs and FNRs derived from MAX-DOAS observations are less well correlated (R=0.48 and 0.59, respectively). These observations also showed differing diurnal cycles between near surface mixing ratios and vertical column densities due to variability in the vertical profile that will be increasingly critical to understand given the ongoing shift from sun synchronous to geostationary satellite observations. Using ground-based measurements, we determined FNRs using both surface mole fractions and vertical column densities, finding FNRs derived from surface mole fractions are generally lower than those derived from column based measurements. Evaluating ozone exceedance probability as a function of FNR for both quantities suggests the transition between a VOC limited and NOx limited regimes may begin at lower FNR values than those derived from satellite based measurements in East LA. We find these differences in FNR derived form ground based and satellite based measurements are driven by variability in the vertical distribution of HCHO. These impacts are most pronounced in late afternoon, when ozone exceedances are most prevalent.

#### 1 Introduction

While the atmospheric chemistry over Los Angeles has been extensively studied with both long term measurements and dedicated intensive field campaigns (e.g. Hering and Blumenthal, 1989; Jacob et al., 2010; Ryerson et al., 2013; Nussbaumer et al., 2023) ozone mole fractions in excess of United States ambient air quality standard (NAAQS) of 70 nmol mol<sup>-1</sup> remain a frequent issue across the region, particularly at warmer temperatures (Pusede et al., 2015), pointing to a continued need for measurements of ozone and its precursors in Southern California.

Ozone production is dependent in a non-linear fashion on both nitrogen oxides ( $NO_x \equiv NO + NO_2$ ) and volatile organic compounds (VOCs) (Haagen-Smit, 1952; Sillman et al., 1990).  $NO_x$  is emitted via both natural sources (e.g. wildfires, lightning) and anthropogenic sources, primarily via combustion. VOCs have a variety of sources including biogenic emissions, wildfires,

and anthropogenic emissions. Formaldehyde is often used in studies as a proxy for VOCs due to it being a common product of VOC oxidation reactions (Millet et al., 2006) and ability to be detected with spectroscopic and remote sensing techniques (e.g. Duncan et al., 2010).

Satellite based measurements of ozone precursors that absorb strongly in the ultraviolet such as NO<sub>2</sub> and HCHO have been used to understand ozone production chemistry over wide regions, particularly those without robust surface monitoring networks (e.g. Martin et al., 2004; Duncan et al., 2010; Jin et al., 2017, 2020; Souri et al., 2020). However, the relationship between the retrieved vertical column densities and near surface mixing ratios more relevant to human health is not straight forward and complicates the use of these data to inform interventions to improve regional air quality (e.g. Schroeder et al., 2017). In addition to direct measurements of NO2 and HCHO, ratios of HCHO to NO2 (FNR) obtained via remote sensing have been explored as a potential diagnostic tool for whether ozone formation chemistry is  $NO_x$  limited or  $NO_x$  saturated (e.g. Martin et al., 2004; Duncan et al., 2010; Jin et al., 2017, 2020; Souri et al., 2020). The use of these column measurements and derived ratios to examine near-surface ozone formation chemistry is complicated by variations in the trace gas vertical profile that confound the relationship between the satellite observed vertical column densities and ground level pollutant concentrations which are the target of policy efforts to improve air quality. In particular, results from the DISCOVER-AQ campaign showed that variations in the HCHO and NO<sub>2</sub> vertical profiles lead to ratios of of HCHO/NO<sub>2</sub> columns that do not reflect ozone production conditions at ground level (Schroeder et al., 2017). Inter-comparisons with ground based measurements to try and improve our understanding of these impacts to date have only been undertaken at times centered around a once daily satellite overpass. The ongoing shift from sun synchronous to geostationary satellite observations will require a better understanding of the diurnal behavior of vertical profiles of these trace gases to interpret satellite based column measurements.

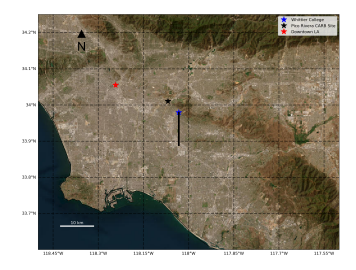
Multiple Axis Differential Absorption Spectroscopy (MAX-DOAS) is frequently used to provide information on both near surface and total column amounts of HCHO and NO<sub>2</sub> concurrently with high temporal resolution during daylight hours and is well suited to provide observational data on how the relationship between these two quantities varies throughout the day. In this work we present two years of vertically resolved measurements of NO<sub>2</sub> and HCHO measurements in southeastern Los Angeles county and their relationship with local ozone conditions. These measurements are the first vertically resolved characterization of these two ozone precursors in southeastern Los Angeles county, improving our understanding of the diurnal variation in ozone precursors and resulting ozone pollution in this part of Southern California.

#### 2 Methods

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#### 2.1 MAX-DOAS Measurements

Multiple axis differential optical absorption spectroscopy (MAX-DOAS) (Hönninger et al., 2004) measurements were made using an Airyx SkySpec-Compact-150 system on the Whittier College campus from 26 March 2020 to 26 May 2022. Whittier College is located approximately 15 km southeast of downtown Los Angeles directly west of the Puente Hills (Fig. 1). The view azimuth of the instrument was 180 degrees east of north, and is shown in Fig. 1 with a black line. Solar spectra were obtained at 1,2,3,5,10,20,40, and 90 degree elevation angles, with one elevation scan taking place over approximately 10 minutes.

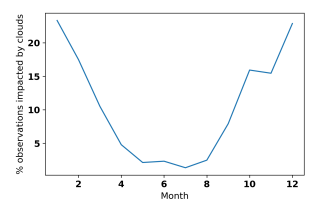


**Figure 1.** Map of the area surrounding Whittier College (Esri, 2024). The location of Whittier College, where the MAX-DOAS measurements took place, is indicated with a blue star. The black line extending from the blue star indicates the view azimuth and approximate horizontal path length of the DOAS observations. The California Air Resources Board site with in situ measurements used in this study is indicated with a black star.

**Table 1.** Median dSCD fitting errors compared to data from standard instruments deployed during the CINDI-2 campaign (Kreher et al., 2020).

Data Product	This study	Standard Instruments (Kreher et al. (2020))
NO <sub>2</sub> (UV)	$8.6 \times 10^{14} \text{ (molec cm}^{-2}\text{)}$	$1\times10^{15}~(\mathrm{molec~cm^{-2}})$
НСНО	$1.9 \times 10^{15}  (\text{molec cm}^{-2})$	8×10 <sup>15</sup> (molec cm <sup>-2</sup> )
O <sub>4</sub> (UV)	$4.1 \times 10^{41} \text{ (molec}^2 \text{ cm}^{-5}\text{)}$	$8 \times 10^{41}  (\text{molec}^2  \text{cm}^{-5})$

DOAS fitting for  $O_4$ ,  $NO_2$ , and HCHO was undertaken using QDOAS (Fayt et al., 2011) following the recommendations from the CINDI-2 intercomparison campaign outlined in Kreher et al. (2020) (Fit windows:  $NO_2$ ,  $O_4$ : 338–370 nm, HCHO: 324.5–359 nm), with exception of the use of a zenith reference spectrum for each elevation scan to minimize the influence of stratospheric trace gases on the spectral fitting. Fits with an RMS value less than 0.001 were not included in further analysis. Typical differential slant column uncertainties, characterized by median fitting errors, shown in Table 1. Cloud screening of these data was done based on the measured colour index ( $\frac{I_{330}}{I_{404}}$ ) using methods detailed in Wagner et al. (2016). This cloud screening removed 8.1% of the data set overall and had a distinct seasonal cycle, shown in Fig. 2, resulting in a dataset that is overly weighted toward spring and summer observations when clear skies were more prevalent. The remaining cloud screened measurements were then used to retrieve vertical profiles of  $NO_2$  and HCHO concentrations averaged over 100 m layers up to an altitude of 4 km, providing vertical profile information for  $NO_2$  and HCHO with 30 min temporal resolution.



**Figure 2.** The percentage of data removed by cloud screening as a function of month. This figure illustrates the larger impact of cloud screening on observations during the winter months.

The retrieval of trace gas vertical profiles from MAX-DOAS slant column densities was performed using HEIPRO (Yilmaz, 2012). The retrieval is a two step process completed using optimal estimation (Rodgers, 2000; Wagner et al., 2004; Frieß et al., 2006). First, the aerosol particle extinction profile is retrieved using slant column density measurements of O<sub>4</sub>, which has a known vertical profile dependent only on temperature and pressure, with the radiative transfer model SCIATRAN (Rozanov et al., 2005) serving as the forward model. For the aerosol extinction profile retrieval, the a priori consisted of an exponentially decaying profile with a scale height of 0.5 km and an aerosol optical depth (AOD) determined by the median measured AOD at the CalTech AERONET site over the measurement period. Similarly, the angstrom exponent (1.23), and phase function (0.70) were also determined using median measured values from the CalTech AERONET site. The surface albedo was assumed to be 0.1 and the aerosol single scattering albedo was assumed to be 0.92 based on measurements during CALNEX (Thompson et al., 2012).

The trace gas retrieval is done similarly using the retrieved aerosol extinction profile as input for the forward model. For  $NO_2$ , the a priori was an exponentially decaying profile with a 0.5 km scale height and a value in the lowest layer of 14 nmol mol<sup>-1</sup>. This value was based on the median of in situ  $NO_2$  measurements during this time period at the California Air Resources Board site in Pico Rivera, 5 km north of the college. For  $CH_2O$ , the a priori was an exponentially decaying profile with a 500 m scale height and a value in the lowest layer of 6 nmol mol<sup>-1</sup> based on measurements during the 2010 CalNex campaign (Warneke et al., 2011).

While the trace gas profiles are typically retrieved on a high vertical resolution altitude grid, in this case every 100 m up to 4 km, this is an over-representation of the actual information content of the original measurements with 40 parameters being retrieved from 7 measurements. Thus, it is desirable to reduce the retrieved profiles to more robust quantities that accurately reflect the information content of the measurements as well as facilitate analysis of long term measurements (Payne et al., 2009; Peterson et al., 2015). With ground-based MAX-DOAS, which is most sensitive near the surface, we reduced the retrieved profiles to a near-surface mole fractions and a lower tropospheric vertical column density (LT-VCD) to emphasize that the

retrieved columns do not necessarily reflect the full tropospheric column (Peterson et al., 2015). Over the course of the study, 93% of  $NO_2$  measurements and 94% of HCHO measurements had more than two degrees of freedom, as determined from the retrieval averaging kernel matrix, indicating that both these quantities could be reliably retrieved from the dSCD measurements. To characterize variability in the vertical distribution of the trace gas of interest, we also calculated the fraction of the retrieved column in the lowest 200 m, which we refer to as the  $f_{200}$  (Peterson et al., 2015). The cloud screened data set consists of 5,790 30 min observations of near-surface mole fractions, LT-VCDs, and  $f_{200}$ s for both  $NO_2$  and HCHO.

#### 2.2 Complementary Data

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Hourly O<sub>3</sub>, NO<sub>2</sub>, and temperature measurements were obtained from the Pico Rivera # 2 South Coast Air Quality Measurement District (SCAQMD) site 5 km northeast of the college (Fig. 1). SCAQMD utilizes Teledyne Advance Pollution Instrumentation, Inc Model T400 instruments to measure hourly ozone at a variety of sites across southern California and makes the date publicly available (http://www.arb.ca.gov/aqmis2/aqmis2.php). NO<sub>2</sub> measurements were made using a Teledyne T200 and also publicly available. Meteorological measurements at SCAQMD sites were made using RM Young Model 81000 Sonic Anemometers. TROPOMI level 2 troposheric NO<sub>2</sub> (Copernicus Sentinel-5P (processed by ESA), 2021) columns and HCHO (European Space Agency, 2020) columns were obtained from the Goddard Earth Sciences Data and Information Services Center (GES DISC). The level 2 TROPOMI swath data were utilized at their native spatial resolution of 5.5x3.5 km without regridding for both species. For both species, only TROPOMI data with a Quality Assurance value greater than 0.75 were used in this analysis.

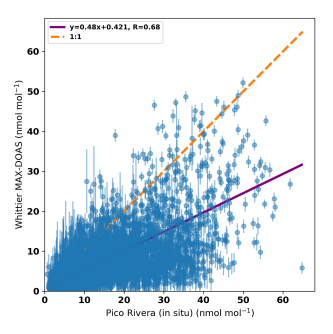
#### 3 Results and Discussion

#### 3.1 Comparisons with in situ data

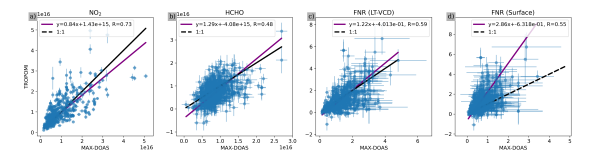
To evaluate the effectiveness of the MAX-DOAS NO<sub>2</sub> retrievals, the mole fractions retrieved for the lowest 100m were correlated with in-situ measurements made by the California Air Resources Board approximately 5 km from the MAX-DOAS. Given the spatial heterogeneity of NO<sub>2</sub>, the comparison of in-situ data to MAX-DOAS retrieved values, which are both horizontally and vertically averaged, should be interpreted with caution. Nevertheless, Figure 3 shows the two measurements are generally well correlated (R=0.68) with the MAX-DOAS retrievals, which are averaged over 100 m vertically and on the order of 10 km horizontally, generally being lower than the in situ measurements at the surface.

#### 115 3.2 Comparisons with TROPOMI

To facilitate comparisons of the ground and satellite-based measurements, MAX-DOAS measurements were averaged over the hour surrounding the local overpass time (~13:30 local time). To compensate for the MAX-DOAS measurements being path-averaged over multiple TROPOMI pixels, the TROPOMI pixel over Whittier and its nearest neighbors were eight directly adjacent pixels (nearest neighbors) were averaged. The horizontal averaging of the MAX-DOAS varies over the study period due to its dependence on aerosol particle extinction in the lower troposphere. For this work, we assume a horizontal path length



**Figure 3.** Correlation of hourly averaged MAX-DOAS measurements with in situ NO<sub>2</sub> data from the California Air Resources Board site in Pico Rivera, 5 km north of the college. The purple line shows the linear regression, while the orange line shows the 1:1 relationship. Error bars for the MAX-DOAS retrievals reflect propagated dSCD measurement errors and a priori error calculated using HEIPRO.



**Figure 4.** a) Correlation of hourly averaged MAX-DOAS NO<sub>2</sub> retrievals with tropospheric columns retrieved from TROPOMI. b) Correlation of hourly averaged MAX-DOAS HCHO retrievals with tropospheric columns retrieved from TROPOMI. c) Correlation of hourly averaged MAX-DOAS LT-VCD based FNRs with those calculated using tropospheric columns retrieved from TROPOMI. d) Correlation of hourly averaged MAX-DOAS surface based FNRs with those calculated using tropospheric columns retrieved from TROPOMI. The purple lines show the orthogonal distance regression, while the black dashed lines shows the 1:1 relationship. Uncertainties for both retrievals are shown in blue. Error bars for the MAX-DOAS retrievals reflect propagated dSCD measurement errors and a priori error calculated using HEIPRO. Error bars for the TROPOMI observations reflect the spatial variability over the nearest neighbor pixels as calculated by the standard error of the mean.

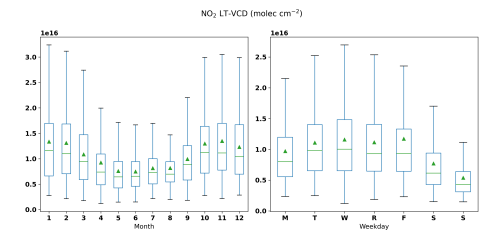
of  $\sim 10$  km based on work by Irie et al. (2011). Error bars for the MAX-DOAS retrievals reflect propagated dSCD measurement errors and a priori error calculated using HEIPRO. Error bars for the TROPOMI observations reflect the spatial variability over the nearest neighbor pixels as calculated by the standard error of the mean. The resulting comparisons for both NO<sub>2</sub> and HCHO over the study period are shown in Fig. 4. The daily NO<sub>2</sub> measurements are generally well correlated (R=0.73), and monthly averages are even more well correlated (R=0.89), although the TROPOMI tropospheric column measurements are generally 16% lower than the ground-based NO<sub>2</sub> column measurements over this study. These results are consistent with other intercomparisons of satellite and ground-based measurements (e.g. Jin et al., 2016; Wang et al., 2017; Pinardi et al., 2020; Chan et al., 2020; Ryan et al., 2023).

Formaldehyde measurements are less well correlated (R=0.48), although monthly averages are still well correlated (R=0.82). The comparison of this finding with prior comparisons is more muddled than that of NO<sub>2</sub>. TROPOMI HCHO columns retrieved are generally higher than those retrieved via MAX-DOAS over the course of the study. This discrepancy is in contrast to findings from some prior global studies (e.g. Chan et al., 2020; De Smedt et al., 2021), but are consistent with inter-comparisons done in polluted urban areas like London (Ryan et al., 2023) and Kinshasa (Yombo Phaka et al., 2023). Additionally, Whittier is impacted by sea breeze circulation which can introduce errors on the order of 20-30% in the air mass factors used to retrieve HCHO VCDs (Souri et al., 2023b). There are times when TROPOMI retrieves large column densities that are not supported by ground based measurements. These events typically occur in the summer months and are potentially due to impacts of long range transport of wildfire plumes further aloft, which can complicate the retrieval of HCHO from satellite-based measurements (Zhao et al., 2022). It is also important to note that ground based MAX-DOAS retrievals, being most sensitive in the lowest 1-2 km of the atmosphere, do not represent the full tropospheric column which would be retrieved from a satellite based measurement. This distinction is more relevant in the measurement of something produced throughout the troposphere like HCHO than something like NO<sub>2</sub> which is primarily emitted near the surface.

#### 3.3 Diurnal variations

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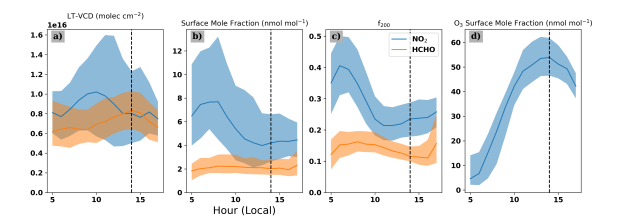
Figure 6 shows diurnal cycles of the vertical distribution of both NO<sub>2</sub> and HCHO retrieved from MAX-DOAS observations. NO<sub>2</sub> vertical columns ranged between 1.2e15-2.9e17 molecules cm<sup>-2</sup> with a median LT-VCD of 7.9e15. Near surface mole fractions ranged from near zero to 52 nmol mol<sup>-1</sup> with a median value of 4.8 nmol mol<sup>-1</sup>. NO<sub>2</sub> vertical distributions tend to be more surface based events have more of the column in the lowest 200 m of the atmosphere in the morning with peak near surface mole fractions occurring between 6 and 8 am local time (Fig. 6b) and peak LT-VCDs occurring closer to noon (Fig. 6a). The observed peak near surface mole fractions and LT-VCDs are generally lower than those seen in Kim et al. (2016b) during the 2010 CalNex campaign at all times of day. This discrepancy could also reflect the fact that our observations include weekend observations which have lower NO<sub>2</sub> throughout the study period (Fig. 5), seasonal differences since CalNex took place during the summer and our observations encompass all seasons, ongoing local NOx emission reductions (Duncan et al., 2016), or impacts of path averaging on MAX-DOAS retrievals. While CalNex took place in the summer and our data set spans all seasons, this is not a likely explanation as summer NO<sub>2</sub> LT-VCDs were typically lower than other months over the course of this study (Fig. 5). It should also be noted that our study period includes a time period encompassing COVID-19 related



**Figure 5.** Box and whisker plots show the seasonal cycle for NO<sub>2</sub> LT-VCDs (left) and clear weekend effect (right). For both plots, the box encompasses the inner quartiles, the green line shows the median, the green triangle shows the mean, and the whiskers show the 5th to 95th percentile range.

lockdown periods which likely altered local NOx emissions. While our data set does not include observations prior to the lockdown period, work by Goldberg et al. (2020) shows TROPOMI NO<sub>2</sub> columns decreased by 32.6% in the Los Angeles area from March 15th 2020 to April 30th 2020 after accounting for differences in meteorology and solar geometry. Our observed diurnal cycle shows NO<sub>2</sub> peaking for both quantities earlier in the day than observed by Kim et al. (2016b), potentially due to differing local sources and traffic patterns in Whittier compared to Pasadena. The fraction of NO<sub>2</sub> retrieved in the lowest 200 m peaks in the early morning at approximately 40% before decreasing to around 20% in the late morning and staying there through the afternoon (Fig. 6c). The median retrieved f<sub>200</sub> was 26%. These dynamics are consistent with boundary layer dynamics typically observed in Southern California (Rahn and Mitchell, 2016).

In contrast, HCHO columns are consistently more vertically distributed with the fraction of HCHO at the surface being between 10 and 20% (Fig. 6c) with a median of 15%. This finding likely reflects the production of HCHO throughout the boundary layer, rather than just near the surface (Schroeder et al., 2017). Sea breeze circulation, which does impact Whittier, could also explain the vertical distribution of HCHO aloft (Souri et al., 2023b). HCHO LT-VCDs ranged between 1.6e15 to 9.7e16 molecules cm<sup>-2</sup> with a median LT-VCD of 6.4e15. Near surface mole fractions ranged from near zero to 14.9 nmol mol<sup>-1</sup> with a median value of 1.9 nmol mol<sup>-1</sup>. Retrieved near surface mole fractions also do not show any diurnal variability, with the median observations consistently being 2 nmol mol<sup>-1</sup> (Fig. 6b) throughout the day. In contrast, median formaldehyde LT-VCDs increase throughout the day peaking in the early afternoon (Fig. 6c), which has been seen in other HCHO column measurements with biogenic VOC sources(e.g. Ryan et al., 2020). Given the high temperatures and large amount of vegetation around the measurement site, it is likely this peak in HCHO can be at least partially attributed to biogenic sources (e.g. Kaiser et al., 2015; Zhao et al., 2022; Chen et al., 2023). This peak in HCHO also coincides with the daily peak in median ozone mole fractions measured at the nearby California Air Resources Board site in Pico Rivera.



**Figure 6.** Plots show the median observed values (solid line) as well as the inner quartile range (shaded regions) for  $NO_2$ (blue) and HCHO (orange) binned by hour of the observation. From left to right the plots show, lower tropospheric vertical column density (a), near surface mole fraction (b), and the  $f_{200}$ , the fraction of the retrieved column in the lowest 200 m of the atmosphere (c). The median time of peak ozone mole fractions observed at the Pico Rivera CARB site is denoted with a vertical dashed line. Panel d shows the average diurnal ozone cycle over the study period.

#### 175 3.4 Examining Ozone Production Chemistry with MAX-DOAS

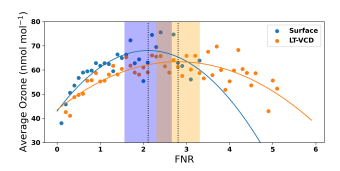
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Ozone formation is dependent on both volatile organic compounds and nitrogen oxides (Sillman et al., 1990). Depending on the relative abundance of both quantities, ozone production can be characterized as NOx limited, where reductions in NOx emissions will be more effective limiting ozone formation, and NOx saturated, where VOC emissions reductions will be more effective limiting ozone formation. Diagnosis of the ozone formation regime requires insights into the relative abundance of both VOCs and NOx. To obtain this information from remote sensing measurements, formaldehyde and NO<sub>2</sub> retrievals can be used (e.g. Martin et al., 2004). FNR values are commonly derived from satellite based remote sensing measurements to provide insights on regional scale ozone production chemistry (e.g. Martin et al., 2004; Duncan et al., 2010; Jin et al., 2017, 2020; Souri et al., 2020). Comparisons of satellite based FNR with surface ozone mole fraction measurements during summer months (May-October) have been used to determine FNR values indicating a transition between NOx limited and NOx saturated ozone production (e.g. Jin et al., 2020).

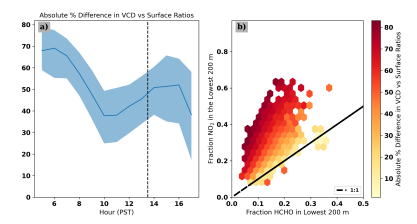
To estimate threshold FNR values from MAX-DOAS measurements, we paired hourly averaged MAX-DOAS observations during all times the temperature exceeded 25° with ozone measurement data from the Pico Rivera CARB site. We then calculated the probability of the observed ozone exceeding the United States ambient air quality standard of 70 nmol mol<sup>-1</sup> as a function of FNR using both near surface mole fractions and LT-VCD, both of which are retrieved concurrently. For each method, the maximum probability of an ozone exceedence as a function of FNR and its associated uncertainty was determined using a 2nd order polynomial fit following methods described in Jin et al. (2020). Figure 7 shows FNR values derived from retrieved surface mole fractions are generally lower than those derived from LT-VCD measurements. This decrease in FNR



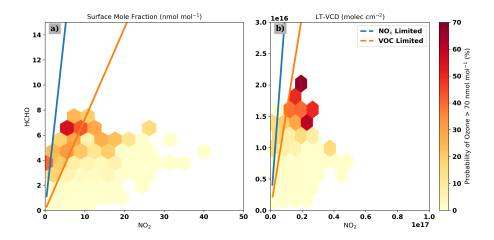
**Figure 7.** Determination of ozone formation regimes from ground-based MAX-DOAS data. Ozone values binned by surface FNR are shown in blue, while those binned by LT-VCD FNR are shown in orange. Second order polynomial fits are shown using solid lines, with the shaded region representing the uncertainty in the maximum value. Dashed vertical lines represent the maximum probability of an ozone excedence as a function of FNR for both methods.

near the surface was also observed in an analysis of MAX-DOAS measurements in three Chinese cities (Hu et al., 2024), which also utilized vertically resolved ozone measurements to show an increasing frequency of NOx limited ozone production as altitudes increased up to 1 km. The transition ranges in this work, determined from near surface FNR and LT-VCDs have less spread than those observed in the Hu et al. (2024) work suggesting that there is less variation in ozone production with altitude observed in Whittier over this time period. The combination of ranges suggests the range of FNR values indicating a transition between NOx limited and NOx saturated ozone production in Whittier is between 1.6 and 3.3 (Fig. 7). Restricting this analysis to the TROPOMI overpass time to facilitate comparisons with other satellite-based FNR measurements showed this transition between FNR values of 0.7 to 3.4. Satellite based estimates of threshold values of this ratio that would indicate a transition in ozone formation chemistry regimes vary regionally but were found to be between 4 and 5 over the Los Angeles metropolitan area based on a study of 20 years of satellite-based FNR observations (Jin et al., 2020). The MAX-DOAS measurements in this study suggest this transition between production regimes occurs at a lower FNR in Whittier than observed over LA generally.

The differences between surface and column based FNR retrievals is most pronounced in the early morning and mid afternoon, reflecting the diurnal variability of the vertical profiles of NO<sub>2</sub> and HCHO (Fig. 8). Near surface ozone exceedances are more likely when HCHO is enhanced (Fig. 9), this leads to a disconnect between satellite based observations since HCHO profiles tend to have the bulk of the HCHO present aloft, away from the surface (Fig. 6).. This variability in HCHO vertical distribution drives the deviation between surface-based and column based FNRs potentially indicating a misdiagnosis of ozone production regimes based on satellite based observations over Whittier as suggested in Schroeder et al. (2017) and reflected in the relatively poor agreement of FNRs between satellite and ground-based observations seen in this study (Fig. 4c). Large uncertainties in HCHO columns retrieved from TROPOMI (Fig. 4) likely also contribute to the poor agreement between TROPOMI and ground-based ratios as seen in Souri et al. (2023a). The interpretations of these ratios is also complicated by the impacts of NO<sub>x</sub> on VOC oxidation which impacts the formation of HCHO and complicates the use of HCHO as a VOC



**Figure 8.** Panel a shows the diurnal variation of the FNR difference expressed as a percentage calculated using surface based quantities vs column based quantities. The solid line represents the median hourly observation and the shaded region represents the inner quartiles. The TROPOMI overpass time is denoted with a vertical dashed line. Panel b shows a hexagonally binned scatter plot of the distribution of FNR differences as a function of vertical distribution of ozone precursor. Each hexagon represents a bin, with color intensity indicating the average difference between the FNR near the surface and throughout the column within that region.



**Figure 9.** Panel a shows shows a hexagonally binned scatter plot of the probability of an ozone observation exceeding 70 nmol  $\text{mol}^{-1}$  as a function of  $\text{NO}_2$  and HCHO near surface mixing ratios. The blue and orange lines delineate NOx limited and NOx saturated regimes derived in Fig. 7. Panel a shows shows a hexagonally binned scatter plot of the probability of an ozone observation exceeding 70 nmol  $\text{mol}^{-1}$  as a function of  $\text{NO}_2$  and HCHO LT-VCDs. The blue and orange lines delineate NOx limited and NOx saturated regimes derived in Fig. 7.

proxy(Chan Miller et al., 2017). During the study period, we observe that increases in ozone exceedance probability are driven by enhancements in HCHO regardless of NO<sub>2</sub> amount (Fig. 9).

#### 4 Conclusions

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For the first time, MAX-DOAS was used to retrieve the vertical distribution of ozone precursors NO2 and HCHO over eastern Los Angeles. The retrieved LT-VCDs for NO2 are well correlated with satellite based measurements from TROPOMI. Retrieved LT-VCDs for HCHO, while less well correlated than NO2 are consistent with prior comparisons, and reinforce the need to improve HCHO retrievals from satellite based measurements. While the vertical distribution of NO<sub>2</sub> reflects increasing boundary layer height throughout the day, HCHO has a much more consistent vertical distribution with 80% of the column being present outside the lowest 200 m regardless of amount. These diurnal variations in both column and vertical profile complicate the interpretation of columns retrieved from geostationary satellite based observations. As an example the observed variability in vertical profiles can lead to differing diagnostics of ozone production via formaldehyde to NO2 ratios. Using MAX-DOAS retrievals, FNRs can be derived from both near surface and column based quantities. Comparison of both methods to observed ozone yields similar transition values from a NOx limited to NOx saturated regimes, but the agreement between the two FNR varies considerably throughout the day, reflecting diurnal variations in both the column and vertical distribution of both precursor species. These findings point to the need to consider the vertical distribution of NO2 and HCHO when interpreting FNRs derived from satellite based measurements to accurately diagnose ozone production chemistry using geostationary satellite observations, particularly in the late afternoon which has larger differences between surface and column based FNR and coincides with times when enhanced near surface ozone mole fractions associated with adverse impacts on human health are most prevalent.

Data availability. MAX-DOAS retrievals of NO<sub>2</sub> and HCHO lower tropospheric column densities and near surface mole fractions are available at https://doi.org/10.5281/zenodo.11117573(Peterson, 2024). Data from the Pico Rivera #2 surface monitoring station used in this paper are available at California Air Resource Board (CARB)'s Air Quality and Meteorological Information System (http://www.arb.ca.gov/aqmis2/aqmis2.php). Differential slant column densities are available from the corresponding author upon request.

Author contributions. PKP led data analysis, drafted the manuscript, and supervised all work done on the manuscript. EH and LT participated in data analysis. AD maintained the MAX-DOAS instrument during this study. All authors reviewed and contributed to the submitted manuscript.

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