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## Impact of high-resolution *a priori* profiles on satellite-based formaldehyde retrievals

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## Abstract

Formaldehyde (HCHO) is either directly emitted from sources or produced during the oxidation of volatile organic compounds in the troposphere. It is possible to infer atmospheric HCHO concentrations using space-based observations, which may be useful for studying emissions and tropospheric chemistry at urban to global scales depending on the quality of the retrievals. In the near future, an unprecedented volume of satellitebased HCHO measurement data will be available from both geostationary and polarorbiting platforms. Therefore, it is essential to develop retrieval methods appropriate for the next-generation satellites that measure at higher spatial and temporal resolution than the current ones. In this study, we examine the importance of fine spatial and temporal resolution a priori profile information on the retrieval by conducting approximately 45,000 radiative transfer model calculations in the Los Angeles Basin megacity. Our analyses suggest that an air mass factor (AMF, a factor converting observed slant columns to vertical columns) based on fine spatial and temporal resolution a priori profiles can better capture the spatial distributions of the enhanced HCHO plumes in an urban area than the nearly constant AMFs used for current operational products. For this urban area, the AMF values are inversely proportional to the magnitude of the HCHO mixing ratios in the boundary layer. Using our optimized model HCHO results in the Los Angeles Basin that mimic the HCHO retrievals from future geostationary satellites, we illustrate the effectiveness of HCHO data from geostationary measurements for understanding and predicting tropospheric ozone and its precursors.

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1. Introduction

Formaldehyde (HCHO) is directly released to the atmosphere from the sources that

include motor vehicles, industrial activities, prescribed burnings, and wildfires. HCHO is

one of the Hazardous Atmospheric Pollutants (HAP) – that are harmful to human health –

defined by the US Environmental Protection Agency (EPA). More importantly, HCHO is

chemically produced during volatile organic compound (VOC) oxidation processes, and

is therefore correlated with major chemical species formed during photochemical smog

episodes [e.g., ozone (O<sub>3</sub>)]. Because of the close relationship between HCHO and its

VOC precursors, the ratio of satellite HCHO columns to nitrogen dioxide (NO<sub>2</sub>) columns

has been suggested as an indicator of photochemical regimes, i.e., the ratio determines

"VOC-limited (or sensitive)" or "NO<sub>x</sub>-limited (or sensitive)" regimes of O<sub>3</sub> formation in

a certain location and season (Martin et al., 2004). In the presence of NO<sub>x</sub>, HCHO can be

a major source of hydroxyl radical (OH), the most important chemical species in the

troposphere initiating photochemical chain reactions. The chemical lifetime of HCHO

with respect to loss by OH reaction and photolysis is several hours (Warneke et al., 2011).

HCHO is highly soluble and may contribute to aqueous chemical processes in clouds and

precipitation in the atmosphere and in bodies of water at the Earth's surface (Barth et al.,

2007.; Luecken et al. 2012).

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Due to its importance to tropospheric chemistry, atmospheric chemists and the environmental remote sensing community have sought to produce high quality tropospheric HCHO retrievals. Because of its weak absorption in the ultraviolet (UV) spectral region, HCHO is regarded as one of the most difficult species to retrieve from satellite-based radiance observations in the UV-visible (UV-VIS) spectral region (e.g., GOME/GOME-2, SCIAMACHY, OMI, and OMPS; see Martin et al., 2003, Zhu et al., 2016 for references). Therefore, it is important to identify factors affecting the accuracy of HCHO retrievals and to find a method to reduce these uncertainties. The largest uncertainties in satellite trace gas retrievals based on UV-VIS spectral measurements arise from the calculation of the air mass factor (AMF), which converts the slant column density of a trace gas to its vertical column values by considering the vertical sensitivity of the observations (AMF = slant column/vertical column, Palmer et al., 2001; Boersma et al., 2004). Palmer et al. (2001) expressed the AMF as a vertical integral of the product of scattering weight functions and normalized vertical profile shapes of trace gases that vary with atmospheric heights. The scattering weight function can be pre-calculated in a look-up table using radiative transfer (RT) model simulations, while the profile shapes are generally derived from a three-dimensional chemical transport model. This formulation has been widely used to derive operational trace gas retrieval products.

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In this study, we examine the role of trace gas vertical profile shapes on HCHO retrievals in the Los Angeles (LA) Basin megacity. The HCHO retrievals from existing polar-orbiting satellites were investigated in previous studies (e.g., Palmer et al., 2001; Millet et al., 2008; Stavrakou et al., 2015; Abad et al., 2015; Zhu et al., 2016); these studies focused on regions with large biogenic sources or used the contrast between land and ocean. Zhu et al. (2014) estimated the anthropogenic VOC emissions from large industrial complexes in Houston, Texas, by oversampling OMI HCHO columns. In the near future, HCHO retrievals will be available from both geostationary [e.g., TEMPO (Fishman et al., 2012; Zoogman et al., 2017), GEMS (Kim et al., 2012), Sentinel-4 (Ingmann et al., 2012, Veihelmann et al., 2015)] and polar-orbiting (e.g., TROPOMI) platforms with much finer temporal and spatial resolutions, enabling satellite-based air quality studies at sub-urban to urban scales. HCHO retrievals at these scales may need a better strategy to deal with spatial and temporal variability in a priori vertical profiles of measured tracers than current methods that rely on profile shapes generated by coarse (horizontal grid resolutions of 2-3 degrees) global models. For example, Heckel et al. (2011) investigated the impacts of the spatial resolution of a priori profiles on NO<sub>2</sub> retrievals in a coastal city (San Francisco, California), which highlighted the need for high resolution a priori data to quantitatively probe tropospheric pollution in coastal

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regions and near localized sources such as power plants. Russell et al. (2011) also found

non-negligible impacts of high spatial and temporal resolution terrain and profile inputs

on the Ozone Monitoring Instrument (OMI) NO<sub>2</sub> retrievals. Kwon et al. (2017)

emphasized the importance of using hourly varying HCHO AMF for geostationary

satellite measurements in East Asia mainly due to temporal changes in aerosol chemical

composition and vertical distributions.

In this study, we simulate fine-resolution (4 km x 4 km) vertical profiles for

HCHO retrievals, and investigate the spatiotemporal variability of the HCHO AMF based

on these profiles. We also show the usefulness of detailed spatial and temporal

information on HCHO plume structures at an urban scale for interpreting the

effectiveness of ozone pollution controls.

2. Data and models

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2.1. Aircraft and ground-based measurements

15- NOAA WP-3 aircraft observations

During the California Nexus of Air Quality and Climate Change (CalNex) campaign, the

NOAA WP-3 aircraft performed 20 research flights mainly over the LA Basin and the

Central Valley in California during May and June 2010 (see Ryerson et al., 2013 for

more information). The main goals of CalNex were to quantify the emissions of

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greenhouse gases and ozone and aerosol precursors and to understand the chemical

transformations and the transport of pollutants. The NOAA WP-3 aircraft was equipped

with a large suite of gas phase and aerosol measurements. In this study, we use the

HCHO measurement of a Proton-Transfer-Reaction Mass-Spectrometry (PTR-MS)

instrument onboard the WP-3 aircraft (Warneke et al., 2011). Airborne HCHO

measurements by PTR-MS are difficult due to a strong humidity dependency. The

detection limit for HCHO with this instrument is between 100 pptv in the dry free

troposphere and 300 pptv in the humid marine boundary layer. The PTR-MS HCHO

measurements have been shown to agree with Differential Optical Absorption

Spectroscopy (DOAS) observations (Stutz and Platt, 1997; Platt and Stutz, 2008) within

the stated uncertainties.

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UCLA long-path DOAS data in Pasadena during CalNex

UCLA's long-path (LP) DOAS instrument (Stutz and Platt, 1997; Platt and Stutz, 2008)

is located on the California Institute of Technology campus on the roof of the Millikan

Library at 35 m AGL (above ground level). Four retro-reflectors are situated northeast of

the main instrument in the mountains behind Altadena at 78, 121, 255, and 556 m AGL.

The average distance between the LP-DOAS telescope and the reflectors is about 6 km.

Spectral retrievals of HCHO mixing ratios were performed in the 324-346 nm

wavelength range using a combination of a linear and non-linear least squares fit, as

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described in Stutz and Platt (1996) and Alicke et al. (2002). Spectral absorption features

of O<sub>3</sub>, NO<sub>2</sub>, HONO, O<sub>4</sub>, and HCHO were incorporated in the fitting procedure. The

campaign-averaged statistical HCHO error in the DOAS measurements during CalNex

was about 150 pptv (Warneke et al, 2011). In the present study, we use these ground-

based DOAS data since vertical distribution information resulting from the four

retroreflectors at different altitudes allows for comparison with the model results.

2.2. WRF-Chem model

We use version 3.4.1 of the Weather Research and Forecasting-Chemistry model (WRF-

Chem, Grell et al., 2005). The model physical and chemical settings are the same as that

used by Kim et al. (2016). The mother and the nested domains of the WRF-Chem model

are the western U.S. (12 km x 12 km horizontal resolution) and the state of California (4

km x 4 km horizontal resolution), respectively. The model has 60 vertical levels with ~50

m thickness between vertical levels up to 4 km above ground level, with coarser vertical

resolution at higher levels. The first model level where mixing ratios of chemical species

are calculated is ~25 m. The simulation period is 26 April 2010 - 17 July 2010.

Meteorological initial and boundary conditions are based on NCEP Global Forecast

System data. The MOZART (Model for OZone And Related chemical Tracers,

http://www.acom.ucar.edu/wrf-chem/mozart.shtml) (Emmons et al., 2010) global model

results are used as initial and boundary conditions for the mother domain of WRF-Chem.

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Biogenic emissions are based on the Biogenic Emissions Inventory System (BEIS)

version 3.13, with additional emissions from urban vegetation (Scott and Benjamin,

2003) are added. The Noah land surface model, YSU planetary boundary layer model,

Lin microphysics scheme, and Grell-Devenyi ensemble cumulus parameterization (only

for the mother domain) are adopted (see references in Kim et al., 2009). The chemical

mechanism is based on the Regional Atmospheric Chemistry Mechanism (RACM)

(Stockwell et al., 1997) with ~30 reaction rate coefficients updated (Kim et al., 2009).

We adopt the NO<sub>x</sub> and CO emission estimates from Kim et al. (2016) that utilized

the fuel-based approaches of McDonald et al. (2012; 2013; 2014). For VOC emissions,

we used the emission estimates from the top-down approach employing ground-based

observations in Pasadena, as described by Borbon et al. (2012), along with the US EPA

NEI05 (US EPA 2008; Kim et al., 2011; 2016) and NEI11 (US EPA 2015; Ahmadov et

al., 2015) inventories. The HCHO model results using the top-down VOC emissions

approach are the focus of this manuscript.

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2.3. VLIDORT radiative transfer model

We used the Vector Linearized Discrete Ordinate Radiative Transfer (VLIDORT) model

(Spurr, 2008) to calculate a trace gas AMF by vertically integrating the product of the

scattering weight function and the normalized vertical profile function of the trace gas, as

described by Palmer et al. (2001). VLIDORT is a multiple-scattering discrete ordinates 20

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RT model for stratified atmospheres. It applies the pseudo-spherical approximation to

solve for the multiple scattering of photons in a stratified atmosphere; diffuse scattering is

evaluated in a plane-parallel medium, but solar attenuation is performed in a spherical

atmosphere. Solar photon single scattering and viewing paths are treated precisely in a

spherically curved atmosphere. Atmospheric and surface thermal emission treatments are

based on ingestion of Planck functions specified at the Earth's surface and at atmospheric

layer boundaries. Since VLIDORT is linearized, simultaneous generation of any number

of analytically derived Jacobians with respect to profile quantities, column quantities, or

surface properties is possible. We adopt the spectral resolution of 0.2 nm and a spectral

range of 300.5-365.5 nm for our HCHO retrievals. Vertical profiles of HCHO, O<sub>3</sub>, NO<sub>2</sub>,

SO<sub>2</sub>, and BrO mixing ratios were used as inputs to the VLIDORT simulations. We used

the WRF-Chem model described above to generate profiles of HCHO, O<sub>3</sub>, NO<sub>2</sub> and SO<sub>2</sub>,

while for BrO, GEOS-Chem global model results were utilized.

3. Results

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3.1. Observed and simulated HCHO

In order to use the model HCHO profiles for AMF calculations and to explore impacts of

fine-resolution a priori on the retrievals, they should be reasonably good representations

of the real atmospheric profiles. Therefore, we evaluate WRF-Chem HCHO simulations

with the ground-based LP-DOAS data and aircraft PTR-MS observations. Figure 1 shows 20

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diurnal variations of the near-surface LP-DOAS HCHO observations and model results

using various emission inventories on weekdays and weekends. The model results using

either the top-down VOC emission estimates based on Borbon et al. (2012; red lines) and

the NEI05 (Kim et al., 2016; blue lines) agree with the observations best. The model

underestimates the LP-DOAS HCHO observations when we ignore the biogenic VOC

emissions or adopt the most-up-to-date VOC inventory for year 2010 (NEI11, described

in Ahmadov et al., 2015), with its lower anthropogenic alkene emissions than those from

the NEI05 and top-down approaches. Maximum observed and modeled HCHO mixing

ratios in Pasadena are about 4 ppbv during weekdays or 5 ppbv during weekends. During

the weekends, faster photochemistry due to lower NO<sub>x</sub> emissions causes higher ozone

and HCHO mixing ratios (Pollack et al., 2012; Kim et al., 2016).

Figure 2 shows the vertical profiles of potential temperature and HCHO mixing

ratio from the aircraft observations and model results in the LA Basin on May 4, 2010.

The potential temperature profiles in the model agree with the observations and help to

characterize different vertical mixing regimes: a stable boundary layer near Catalina

Island and the growth of the convective boundary layer from the LA urban cores eastward

to the desert on the east side of the Basin. Similarly, the WRF-Chem HCHO profiles are

in good agreement with the WP-3 PTR-MS observations. The convective boundary layer

develops mainly by buoyancy forcing during daytime and leads to well-defined boundary

layer heights (or mixing heights) ranging from a few hundred meters to several

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kilometers and well-mixed vertical profiles of potential temperature and scalars.

Meanwhile, stable boundary layers are characterized by a shallow boundary layer

(boundary layer height of maximum a few hundred meters), a positive vertical gradient of

potential temperature near the surface, and poorly-mixed vertical profiles of scalars

because of weak turbulent mixing that frequently occurs over the ocean or during

nighttime. Overall, our model results agree with the observations from the aircraft and

ground-based observations; therefore, it is reasonable to use the model HCHO profiles as

inputs to VLIDORT and to examine the AMF results from this RT model.

3.2. Spatial distribution of AMF and sensitivity to a priori profiles at different times

of day

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The spatial distribution of the VLIDORT HCHO AMF using the WRF-Chem profiles at 4

km x 4 km resolution at different times of day on May 4, 2010 is shown in Figure 3. The

AMF ranges from 0.6 to 1.2 within the LA Basin and in the nearby coastal areas. The

AMF values are 0.6-0.7 in the urban cores. In contrast, for high mountains such as the

Los Padres Forest located in the northwestern part of the Basin, the AMF is greater than 1.

Above the Pacific Ocean near the coast, the AMF is about 0.9-1. These results are similar

to the AMF calculations by Palmer et al. (2001); they obtained AMF = 0.71 in Tennessee,

where high isoprene levels are seen in the boundary layer, and AMF = 1.1 over the North

20 Pacific. The AMF values calculated by Palmer et al. (2001) resulted in Global Ozone

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Monitoring Experiment (GOME) measurements that were ~35% less sensitive to the

HCHO column over Tennessee than over the North Pacific. Palmer et al. (2001) also

noted small AMF values over California, which they attributed to a shallow boundary

layer resulting from strong subtropical subsidence combined with a strong surface source

of HCHO from biogenic hydrocarbons. Our study agrees with this finding, except that

both anthropogenic and biogenic VOC contribute to high formaldehyde in the LA Basin

(Figure 1). General features of the AMF distribution in the area do not change

significantly when a constant surface pressure is used in the RT simulations (see

Supplementary Material Figure S1). The direct influence of complex terrain height on the

AMF is small. Similarly, the spatial pattern was not strongly affected by the emission

inventory used to generate the WRF-Chem HCHO profiles in our study (see

Supplementary Material Figure S1).

Geostationary satellites such as TEMPO (Fishman et al., 2012; Zoogman et al.,

2017), GEMS (Kim et al., 2012), and Sentinel-4 (Ingmann et al., 2012; Veihelmann et al.,

2015) are expected to provide diurnally varying information about tropospheric pollution

during daytime. It is, therefore, useful to investigate if diurnally varying a priori profile

information is needed for accurate retrievals of satellite-based HCHO columns. Figure 3

shows the spatial distribution of VLIDORT HCHO AMF using the WRF-Chem profiles

at 16, 19, and 22 UTC (equivalent to 9, 12, 15 Pacific Daylight Time, PDT, respectively)

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and HCHO columns. Overall, similar patterns of the AMF distribution are shown at all

times: low AMFs in the urban cores and high AMFs in the area of Los Padres National

Forest located in the northwestern region of the Basin. However, there are noticeable

diurnal changes in the AMFs over the high terrain east and northeast of downtown LA

and over the Pacific Ocean near the coast, due to changing photochemical production and

destruction and transport of HCHO throughout the day (Figure 3). Overall, minimum

AMF values are reduced between morning and afternoon as HCHO is photochemically

produced. At 15 PDT, AMF values < 0.6 (the white shading in Figure 3) occur in the

mountainous regions, including the San Gabriel Mountains, San Bernardino National

Forest, Mt. San Jacinto, and Anza-Borrego Desert State Park. Onshore transport of

photochemically produced HCHO plumes from downtown LA to the mountains occurs in

the afternoon (see HCHO columns in Figure 3).

Figure 4 shows vertical distributions of the model HCHO mixing ratios at several

locations in the LA Basin and the Pacific Ocean for the AMF values at different times of

day. Over the Pacific Ocean, the HCHO mixing ratio is small near the surface and more

abundant at higher altitudes. The AMF over the ocean increases with time from 0.86 at 09

PDT to 1.03 at 15 PDT as the HCHO mixing ratio decreases with time, probably due to

transport of the plume from the ocean to the inland area. Over the land, the HCHO

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Padres National Forest where the highest AMF (0.91-1.21) occurs, the boundary layer grows with time, but the mixing ratio of HCHO is small (< 1 ppbv). In Pasadena and at the LA Main St. site, the boundary layer heights and HCHO mixing ratios increase from 9 PDT to 12 PDT. The maximum HCHO value in the boundary layer is about 6 ppbv. The HCHO in the boundary layer decreases at 15 PDT, but mixing ratios above the boundary layer (> 1 km) increase due to the upper level easterly transport of the HCHO plumes. Consequently, the AMF decreases from 0.7 at 9 PDT to 0.6 at 12 PDT and then increases to 0.7 at 15 PDT, due to an enhanced sensitivity to increased upper-level HCHO mixing ratios. For these urban core sites, the HCHO AMF ranges from 0.6 to 0.7. In the San Gabriel Mountains, San Bernardino National Forest, and Mt. San Jacinto, the boundary layer height is well defined and shallow and does not change significantly throughout the day. However, the AMF values change substantially (decreasing by ~40%) throughout the day over these locations; this is likely because HCHO mixing ratios increase between morning and afternoon, mainly due to transport and formation of the plumes originating

mixing ratio is higher in the boundary layer than in the free atmosphere. In the Los

from urban core regions. The AMF at Anza-Borrego Desert State Park decreases with

time from 0.96 at 9 PDT to 0.71 at 15 PDT due to increasing HCHO mixing ratios, in

spite of the increase in boundary layer height. These findings highlight that the

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importance of using time-varying, high spatial resolution a priori profile information for

the accurate retrieval of geostationary HCHO measurements.

We extended this analysis in Figure 5, where for ranges of HCHO AMF (e.g., 1.0

< AMF < 1.1) across the model domain, the model HCHO profiles are averaged and

plotted at the three times (9, 12, 15 PDT). Each plot shows that the AMF values are

smaller when the HCHO mixing ratios are higher near the surface. At 12 and 15 PDT, as

expected, the profiles have more well-mixed shapes for deeper vertical layers. The

dependence of the AMF value on the profile shape is similar at each time of day.

Using all available data points, we investigate the relationship between AMF and

the HCHO mixing ratio at 200 m in the boundary layer at different times of day in Figure

6. The plot illustrates that as the HCHO mixing ratio increases, the AMF decreases. In

general, AMF values decrease from morning to late afternoon.

For UV-VIS retrievals, it is generally assumed that only the vertical profile shape,

rather than the absolute magnitude of the absorber, affects the value of the AMF.

However, our study suggests a strong anti-correlation between the absolute concentration

and the AMF: the AMF is low in the area of intense HCHO plumes. To understand the

importance of the absolute magnitude of HCHO mixing ratios within the context of the

mathematical formula of AMF used, we examine shape factor, scattering weight function,

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and AMF quantitatively. According to Palmer et al. (2001), AMF is expressed as

$$AMF = AMF_G \int_0^\infty w(z) S_z(z) dz.$$
 (1)

Here  $AMF_G$  is a geometric air mass factor that is a function of solar zenith angle and satellite viewing angle, w(z) is a scattering weight that is associated with the sensitivity of the backscattered spectrum to the abundance of the absorber at altitude z, and  $S_z(z)$  is a vertical shape factor for the absorber representing a normalized vertical profile of number density. The vertical shape factor is defined as

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$$S_{z}(z) = \frac{n(z)}{\Omega_{v}} \tag{2}$$

, where n(z) is the number density (molecules cm<sup>-3</sup>) at altitude z and  $\Omega_v$  is the vertical column density or column (molecules cm<sup>-2</sup>) of HCHO. In this manuscript, AMF in Equation (1) is vertically integrated to the top of model domain that is roughly the top of troposphere or above. Therefore, AMF here is tropospheric AMF. To understand the sensitivity of AMF on the vertical distribution, we also define  $\Delta AMF_i$ , a discrete increment of AMF for each model layer.

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$$\Delta AMF_i = AMF_G w_i S_{zi} \Delta z_i \tag{3}$$

, where *i* is an index representing the vertical grids,  $\Delta \mathbf{z_i}$  is the layer depth for the grid *i*, and  $\sum \Delta AMF_i = AMF$ .

In Figure 7, the vertical shape factor in Equation (2), the scattering weight (multiplied by geometric AMF), and  $\Delta AMF_i$  are plotted as a function of height over the North Pacific Ocean, San Gabriel Mountains, and Anza Borrego Desert State Park at 16, 19, and 22 UTC (see Figure 4 for the locations of these sites). The differences in the shape factor over the North Pacific Ocean are clear at altitudes > ~1000 m: the shape factor values at 22 UTC are larger than those at 16 and 19 UTC. In contrast, the HCHO column at 22 UTC is smaller than those at 16 and 19 UTC over the ocean (Figure 4). As the column density value decreases, the shape factor above ~1000 m becomes larger and causes higher  $\Delta AMF_i$  and (tropospheric) AMFs, because a column density value is used as a normalization parameter for a shape factor. In order words, the satellite measurement is more sensitive to the profile at 22 UTC than that at 16 UTC at this point over the Pacific Ocean.

For the San Gabriel Mountain site, the HCHO is confined below ~1400 m at 16, 19, and 22 UTC (there are no significant changes in boundary layer height during this

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time period) and its mixing ratio increases with time (Figure 4). The shape factor at 19

and 22 UTC is higher than that at 16 UTC below ~1400 m altitude (Figure 7, middle row).

However, above this height, the shape factor and  $\Delta AMF_i$  decrease with time: both are

largest at 16 UTC and smallest at 22 UTC. The tropospheric AMF follows  $\Delta AMF_i$ 

above ~1400 m and also decrease with time from 1 to 0.58. Thus, the satellite

measurement is more sensitive to the profile at 16 UTC than that at 22 UTC in this

mountainous area. The plot over the San Gabriel Mountain area illustrates that not only

boundary layer height, but also the absolute magnitude of HCHO, influence the AMF

value.

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Anza-Borrego Desert State Park represents an example of a case in which both

boundary layer height and HCHO mixing ratio increase with time (Figure 4 and Figure 7,

bottom row). In case of the lowest boundary layer height (at 16 UTC), AMF is largest

(AMF=0.98). When the boundary layer height is the highest among the three time periods

(at 22 UTC), the AMF is smallest (AMF=0.71). For Anza Borrego Desert State Park, total

column or near surface HCHO mixing ratio affect the shape factor, which in turn leads to

an AMF that is inversely proportional to the total column or near surface HCHO mixing

ratio. As shown in Figure 7, the shape factor and  $\Delta AMF_i$  above the boundary layer

decrease with time, which causes a decrease in the tropospheric AMF with time.

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In summary, the absolute value of the column or near-surface mixing ratio of

HCHO affects the shape factor as a normalization factor, in particular the value in the free

troposphere (above boundary layer), which dominates the tropospheric AMF. When the

HCHO mixing ratio is low in the boundary layer, the relative importance of the absorber

in the free troposphere increases. Conversely, when the HCHO mixing ratio is high in the

boundary layer, the relative importance of absorber in the free atmosphere decreases. Our

result suggests that a representation of the HCHO AMF using accurate fine-resolution a

priori profile information is critical to identify HCHO plumes and to place better

constraints on VOC emissions.

10 Although the focus of this manuscript is on the shape factor, we also investigate

the impacts of aerosol loading on AMF for the 8 sites shown in Figure 4. When the

aerosol optical properties from the model results are incorporated in our RT model

calculations, the AMF is reduced by  $\sim 10\%$  at the N. Main St. and Pasadena sites and by <

10% at other sites (not shown). Because the model aerosol results were not thoroughly

evaluated and optimized, the analysis of aerosol impact in this study is limited. It is

possible that the simulated aerosol is overestimated, because the emission inventory is

not up to date (e.g., overestimations of black carbon and SO<sub>2</sub> by a least a factor of 3). It is

likely that the actual impact of aerosols on the AMF is relatively small when compared

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with other factors examined here.

3.3. Air quality application of fine-resolution geostationary HCHO columns

In this section, we illustrate the application of future geostationary HCHO retrievals to

the study of air quality, by using the WRF-Chem HCHO columns as a proxy for satellite

data. Figure 8 demonstrates the distribution of the ratio of HCHO to NO<sub>2</sub> tropospheric

vertical columns from the WRF-Chem model in the LA Basin at different times of day

and on weekdays and weekends for May-June 2010. For more information about the

model NO<sub>2</sub> columns, refer to Kim et al. (2016).

Ratios of HCHO to NO<sub>2</sub> columns provide critical information about chemical 10

regimes relevant to controlling ozone pollution (Martin et al., 2004). In Figure 8, the light

blue to blue contours (HCHO/NO2 < 1) represent VOC-sensitive (or VOC-limited) ozone

production regimes, while the pink to the red contours (HCHO/NO<sub>2</sub> > 1) denote NO<sub>x</sub>-

sensitive regimes. During weekdays in 2010, most of the LA Basin is in the VOC-

sensitive regime, where a reduction in NO<sub>x</sub> emissions can cause an increase in O<sub>3</sub>. In the

late afternoon during weekends, the broad polluted area becomes NO<sub>x</sub>-sensitive, so that

NO<sub>x</sub> reductions lead to O<sub>3</sub> decreases.

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Figure 9 shows 2000-2010 trends in surface O<sub>3</sub> from monitors in Pasadena and

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San Bernardino. During this decade, NO<sub>x</sub> emissions were decreasing in the LA Basin,

largely due to better control of motor vehicle pollution (McDonald et al., 2012). On

weekdays during this decade, there was not a declining trend in surface O<sub>3</sub> in Pasadena,

while O<sub>3</sub> increased in San Bernardino. In contrast, on weekends, O<sub>3</sub> decreased between

2000 and 2010 in both Pasadena and San Bernardino. These observed O3 trends are

consistent with analyses of the ratio of HCHO to NO2 columns, and their representation

of VOC/NO<sub>x</sub> sensitivity, shown in Figure 9. Baidar et al. (2015) found that the spatial

extent and the trend of higher O<sub>3</sub> during weekends than weekdays had decreased in the

LA Basin because of the increased tendency of lower O<sub>3</sub> during hot weekends, especially

after the 2008 economic recession. 10

The polar-orbiting satellite instruments that are currently available do not provide

diurnally varying information on HCHO/NO<sub>2</sub> columns and VOC/NO<sub>x</sub> sensitivities,

because these measurements are made once a day in either the morning or early afternoon.

The discussion above makes it clear that future geostationary satellite HCHO and NO<sub>2</sub>

columns will provide useful information about photochemical ozone regimes that could

be used to evaluate pollution control policies.

4. Summary and Conclusions

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Our tests of the sensitivity of HCHO AMF to several factors confirm the importance of a

priori HCHO profile shapes. Our study reveals that, for similar vertical profile shapes,

the absolute magnitude of HCHO concentration is also an essential factor in determining

the AMF. For the coastal LA Basin megacity studied in this work, the AMF values are

inversely proportional to the magnitude of the HCHO mixing ratios in the boundary layer.

Furthermore, the AMF over land is lower in the late afternoon (15 PDT) than in the

morning (09 PDT), because of increasing HCHO mixing ratios throughout the day.

Therefore, diurnal updates and fine spatial resolution a priori profile shapes are likely to

improve the retrievals of satellite-based HCHO columns.

The spatial distributions of fine-scale model HCHO columns in the LA Basin show

hot spots in downtown LA around noon and enhancement and transport of the plumes to

the eastern part of the Basin in the late afternoon. The ratio of HCHO to NO<sub>2</sub> columns

during weekdays and weekends provides information on the chemical regimes relevant to

ozone formation at various locations and times in the Basin. Future geostationary

satellites (e.g., TEMPO) may provide similar information, which could be used to assess

the effectiveness of existing pollution controls and could help in planning or revising air

pollution control policies.

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Lee at Pusan National University.

The WRF-Chem model version 3.4.1 used in this study is available at <a href="http://www2.mmm.ucar.edu/wrf/users/download/get source.html">http://www2.mmm.ucar.edu/wrf/users/download/get source.html</a>. We acknowledge use of MOZART-4 global model output available at <a href="http://www.acom.ucar.edu/wrf-chem/mozart.shtml">http://www.acom.ucar.edu/wrf-chem/mozart.shtml</a>. The CalNex field campaign data are available at <a href="http://www.esrl.noaa.gov/csd/projects/calnex/">http://www.esrl.noaa.gov/csd/projects/calnex/</a>.

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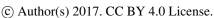
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Atmospheric Chemistry And Physics Discussions

Figure captions

Figure 1. Diurnal variations of ground-based observations (black filled circles with solid

lines) of HCHO and the corresponding model simulations (lines without symbols) in

Pasadena (34.1370°N, 118.1254°W) averaged for weekdays (left) and weekends (right).

All model simulations utilized the fuel-based NO<sub>x</sub> and CO emissions in Kim et al. (2016).

The red solid line shows the results utilizing the VOC emissions from the top-down

approach in Borbon et al. (2012), the red dashed line denotes the same model settings

represented by the red solid line (top-down approach) except for zero biogenic VOC

(BVOC) emissions, the blue solid line represents the model results using the VOC

emissions from NEI05 (as in Kim et al., 2016), and the light blue line shows the model

output using the VOC emissions from NEI11.

Figure 2. The flight path of NOAA WP-3 (top) and the spatial distribution of vertical

profiles of aircraft observed and model simulated potential temperature (middle) and

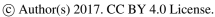
HCHO (bottom) in the LA Basin on May 5, 2010. The black filled circles and red solid

lines/symbols represent the observations and model results, respectively.

Figure 3. The spatial distributions of air mass factors from the radiative transfer model

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calculations (left) and HCHO columns (right) in the LA Basin at 16 UTC (top), 19 UTC

(middle), and 22 UTC (bottom) on May 4, 2010. The black filled circles are included as

points of further investigations, representing background, urban cores, and downwind

regions.

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Figure 4. Vertical profiles of HCHO are shown for various points of interest (red symbols

on a Google map). Blue, orange, and magenta lines represent 16, 19, and 22 UTC,

respectively. The altitude above ground level is shown.

Figure 5. Vertical profiles of HCHO averaged for the AMF value intervals (as in legends)

at 16, 19, and 22 UTC (left to right) are displayed. Thick lines with symbols are for

averages and thin dotted lines are for one standard deviation values. The altitude above

ground level is shown.

Figure 6. The relationship between the AMF and model HCHO volume mixing ratio is

demonstrated. Different colors denote different times. The HCHO mixing ratio at ~200 m

altitude is plotted.

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Figure 7. Vertical profiles of (left) slope factor and scattering weight and (right)  $\Delta AMF_i$ 

(discrete increment of AMF) at North Pacific Ocean, San Gabriel Mountains, and Anza

Borrego Desert State Park. Scattering weights multiplied by geometric AMF are shown.

Figure 8. Spatial distributions of the ratios of the model HCHO column to NO<sub>2</sub> column

during weekdays (left) and weekends (right) at 09 PDT, 12 PDT, and 15 PDT for May-

June 2010. The light pink to red colored contours denote the area under the NO<sub>x</sub>-limited

chemical regime.

10 Figure 9. Decadal O<sub>3</sub> trends in Pasadena and San Bernardino during weekdays (red) and

weekends (blue) are shown. The linear least square fits of O<sub>3</sub> for Wednesday and Sunday

are plotted in dashed lines.

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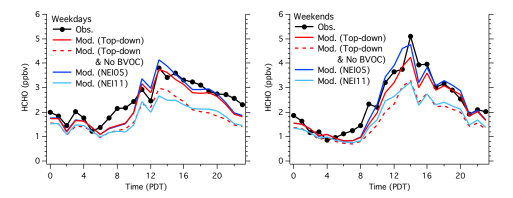


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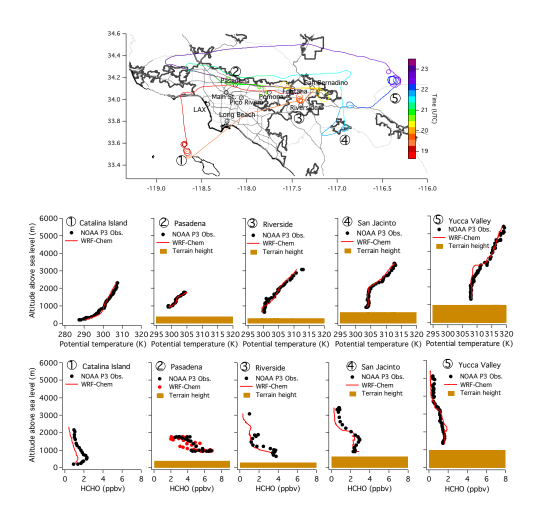


Figure 2. The flight path of the NOAA WP-3 aircraft (top) and the spatial distribution of vertical profiles observed on the aircraft and simulated by the model for potential temperature (middle) and HCHO (bottom) in the LA Basin on May 5, 2010. The black filled circles and red solid lines/symbols represent the observations and model results, respectively.

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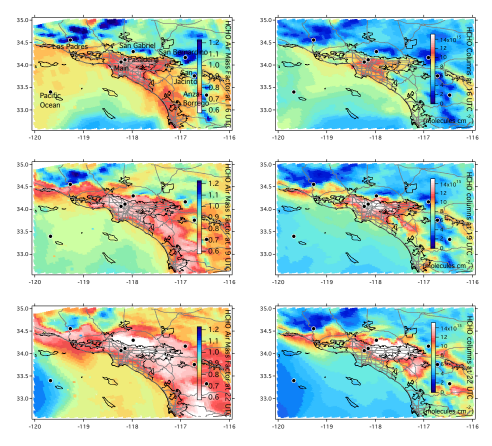


Figure 3. The spatial distributions of air mass factors from the radiative transfer model calculations (left) and HCHO columns (right) in the LA Basin at 16 UTC (top), 19 UTC (middle), and 22 UTC (bottom) on May 4, 2010. The black filled circles are included as points of further investigations, representing background, urban cores, and downwind regions.

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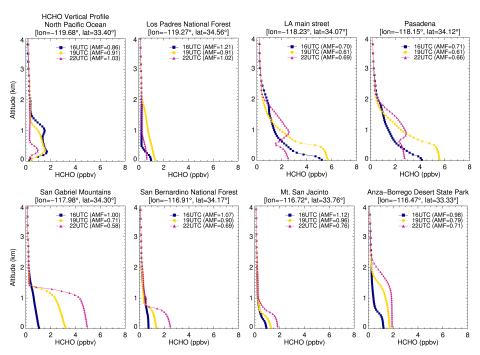


Figure 4. Vertical profiles of HCHO mixing ratio are shown for various points of interest (red symbols on a Google map). Blue, orange, and magenta lines represent 16, 19, and 22 UTC, respectively. The altitude above ground level is shown.

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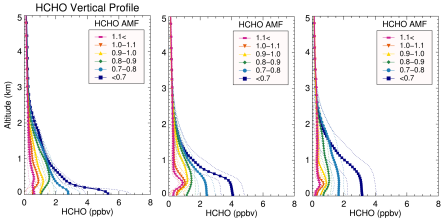


Figure 5. Vertical profiles of HCHO mixing ratio averaged for the AMF value intervals (as in legends) at 16, 19, and 22 UTC (left to right) are displayed. Thick lines with symbols are for averages and thin dotted lines are for one standard deviation values. The altitude above ground level is shown.

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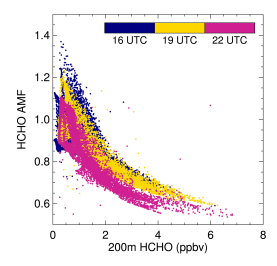


Figure 6. The relationship between the AMF and model HCHO volume mixing ratio is demonstrated. Different colors denote different times. The HCHO mixing ratio at  $\sim$ 200 m altitude is plotted.

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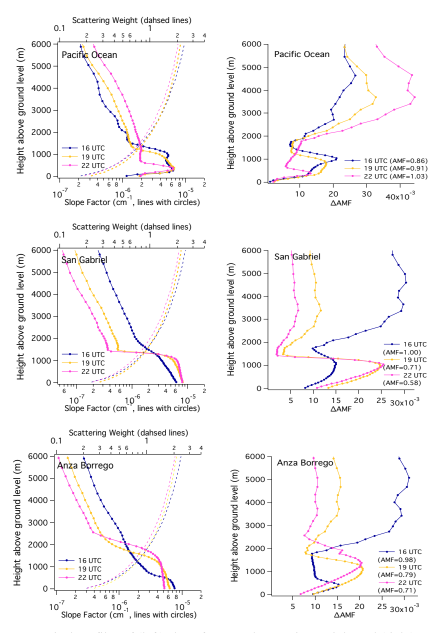


Figure 7. Vertical profiles of (left) slope factor and scattering weight and (right)  $\Delta AMF_i$  (discrete increment of AMF) at North Pacific Ocean, San Gabriel Mountains, and Anza Borrego Desert State Park. Scattering weights multiplied by geometric AMF are shown.

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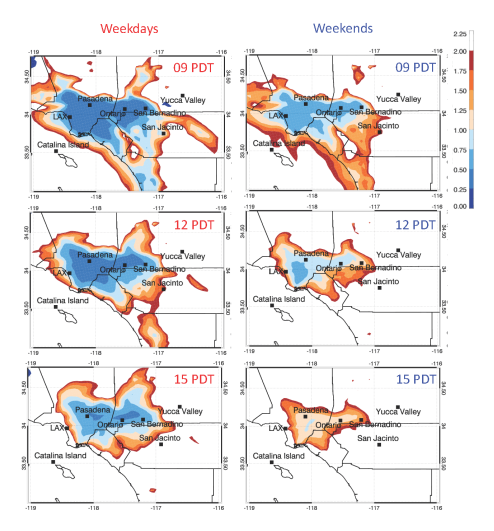


Figure 8. Spatial distributions of the ratios of the model HCHO column to  $NO_2$  column during weekdays (left) and weekends (right) at 09 PDT, 12 PDT, and 15 PDT for May-June 2010. The light pink to red colored contours denote the area under the  $NO_x$ -limited chemical regime, while blue contours denote VOC-limited regions.

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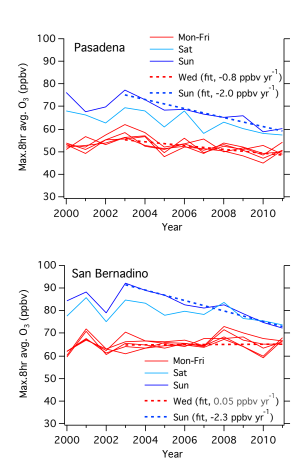


Figure 9. Decadal  $O_3$  trends in Pasadena and San Bernardino during weekdays (red) and weekends (blue) are shown. The linear least square fits of  $O_3$  for Wednesday and Sunday are plotted in dashed lines.