Interannual variability of summertime formaldehyde (HCHO) vertical column density and its main drivers in northern high latitudes

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Abstract:

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The northern high latitudes (50-90°N, mostly including boreal forest and tundra ecosystem) has been undergoing rapid climate and ecological changes over recent decades, leading to significant variations in Volatile Organic Compounds (VOCs) emissions from biogenic and biomass burning sources. Formaldehyde (HCHO) is an indicator of VOC emissions, but the interannual variability of HCHO and its main drivers over the region remain unclear. In this study, we use the GEOS-Chem chemical transport model and satellite retrievals from Ozone Monitoring Instrument (OMI) and Ozone Mapping and Profiler Suite (OMPS), to examine the interannual variability of HCHO vertical column density (VCD) during the summer seasons spanning from 2005 to 2019. Our

results show that in 2005-2019 summers, wildfires contribute 75-90% of the interannual variability of HCHO VCD over Siberia, Alaska, and Northern Canada, while biogenic emissions and background methane oxidation accounts for ~90% of HCHO interannual variability over Eastern Europe. We find that monthly Solar-induced chlorophyll fluorescence (SIF) from Orbiting Carbon Observatory-2 (OCO-2), an efficient proxy for plant photosynthesis, shows a good linear relationship (R=0.6-0.7) with modelled biogenic HCHO column (dVCD_{Bio,GC}) in Eastern Europe, Siberia, Alaska and Northern Canada, indicating the coupling between SIF and biogenic VOC emissions over the four domains on a monthly scale. In Alaska, Siberia and Northern Canada, SIF and dVCD_{Bio,GC} both show a relatively lower interannual variabilities (SIF: CV=1-9%, dVCD_{Bio,GC}: CV=1-2%. CV: Coefficient of Variation) comparing to wildfire-induced HCHO (CV=8-13%), suggesting that the high interannual variabilities of OMI HCHO VCD (CV=10-16%) in these domains are likely driven by wildfires instead of biogenic emissions.

1. Introduction

VOCs are main precursors of tropospheric ozone and secondary organic aerosols, strongly impacting air quality and climate (Atkinson, 2000; Kroll and Seinfeld, 2008; Mao et al., 2018; Zheng et al., 2020). HCHO is mainly produced from atmospheric VOC oxidation with a short photochemical lifetime on the order of hours, serving as an indicator of non-methane VOC (NMVOC) emissions and photochemical processes (Fu et al., 2007; Millet et al., 2008). Understanding the interannual variability of HCHO is important for quantifying long-term trend of VOC emissions in response to climate changes and air quality control implementation.

Several studies suggest that biogenic VOC emissions are largely responsible for interannual variabilities of HCHO on a global scale (Palmer et al., 2001; De Smedt et al., 2008; González Abad et al., 2015; De Smedt et al., 2018). Stavrakou et al. (2009) attributes Biogenic VOCs (BVOCs) emissions as the predominant source of global HCHO columns, in which isoprene alone contributes to 30% of global HCHO. Isoprene emissions were also found to be the major driver of HCHO interannual variability (Bauwens et al., 2016; Stavrakou et al., 2018; Morfopoulos et al., 2022). During wildfire seasons, pyrogenic emission is the secondary important controlling factor of HCHO over the whole Amazon (Zhang et al., 2019) and contributes to 50-72% of HCHO total column in Alaskan summer fire seasons (Zhao et al., 2022). Over Antarctic region, HCHO is produced mainly from methane oxidation with hydroxyl radicals (OH), with possible unknown HCHO sources and long-range transport (Riedel et al., 1999). The interannual variability of HCHO over this region is still unclear.

Northern high latitudes are experiencing rapid Arctic warming in recent decades, resulting in strong increases in BVOC emissions (Lappalainen et al., 2009; Vedel-Petersen et al., 2015; Kramshøj et al., 2016; Seco et al., 2022). Several studies suggest monoterpenes to be the most abundant BVOC species in boreal forests over middle and north Europe, and southeastern Siberia (Spirig et al., 2004; Timkovsky et al., 2010; Bäck et al., 2012; Rantala et al., 2015; Juráň et al., 2017; Zhou et al., 2017). This BVOC speciation appears to be different in the boreal forests in Alaska, Northern Canada and Eastern Siberia, where isoprene appears to be the most abundant BVOC species (Blake et al., 1992; Timkovsky et al., 2010; Zhao et al., 2022). BVOC measurements in tundra systems show a very strong positive temperature dependence for isoprene fluxes, over Greenland (Vedel-Petersen et al., 2015; Kramshøj et al., 2016; Lindwall et al., 2016a),

northern Sweden (Faubert et al., 2010; Tang et al., 2016) and the Alaskan North Slope (Potosnak et al., 2013; Angot et al., 2020; Selimovic et al., 2022).

Wildfire is another important source of HCHO (Permar et al., 2021). A number of studies have shown positive trend and strong interannual variability of wildfires over Arctic regions in the past few decades (Kelly et al., 2013; Giglio et al., 2013; Descals et al., 2022). Several modelling studies suggest that wildfires can become the main source of HCHO over Alaska (Zhao et al., 2022), Siberia and Canada (Stavrakou et al., 2018). In fact, the contribution from wildfires could be even larger as models tend to underestimate the secondary production of HCHO from other VOC precursors (Alvarado et al., 2020; Zhao et al., 2022; Jin et al., 2023). To what extent wildfires contribute to HCHO interannual variability remains unclear.

Solar Induced Fluorescence (SIF) could potentially provide additional constraints on biogenic-related HCHO column over northern high latitudes, due to their similar dependence on temperature and light availability (Foster et al., 2014; Zheng et al., 2015). SIF is the re-emission of light by plants as a result of absorbing solar radiation during photosynthesis and is widely used to estimate vegetation productivity and health (Porcar-Castell et al., 2014; Magney et al., 2019). Isotopic labeling studies show that 70-90% of isoprene production is from chloroplasts, directly linked to photosynthesis (Delwiche and Sharkey, 1993; Karl et al., 2002; Affek and Yakir, 2003). As SIF is directly linked to flux-derived Gross Primary Productivity (GPP) and HCHO can be largely explained by isoprene emissions (Zheng et al., 2017), we expect to use SIF as a valuable tool to constrain biogenic emissions from boreal forest at northern high latitudes.

The new retrievals of HCHO from OMI and OMPS provide a continuous long-term record on a global scale, with improved calibration, updates in spectral fitting and air mass factor calculations (González Abad et al., 2022; Nowlan et al., 2023). Here we use the newly retrieved HCHO vertical column density (VCD) products from OMI and OMPS, combined with GEOS-Chem chemical transport model, to examine summertime HCHO spatiotemporal variability over northern high latitudes from 2005 to 2019. The satellites and the model are introduced in Sect. 2. In Sect. 3, we evaluate the spatial variability of HCHO VCD using satellite retrievals and evaluate BVOC emissions with previous in-situ measurements. In Sect. 4, we evaluate the interannual variability of HCHO VCD using satellite retrievals, and present model sensitivity tests to demonstrate how background HCHO, wildfire and biogenic VOC emissions influence HCHO interannual variability across Alaska, Siberia, Northern Canada and Eastern Europe. In Sect. 5, we evaluate biogenic HCHO interannual variability using satellite SIF data. Summary and discussion are in Sect. 6.

2. Observations and Model

2.1. Observational data sets

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We use satellite observations of tropospheric HCHO columns from OMI and OMPS to evaluate summertime HCHO variability at northern high latitudes. OMI is a UV/Visible backscatter spectrometer on-board the Aura satellite launched in July 2004, with global daily coverage at an overpass time of 13:30 LT. OMI provides a long-term record of HCHO VCD but is discontinued in 2023. OMPS is the continuation of OMI HCHO measurement over polar region. OMPS is a spectrometer on-board two satellites: NASA/NOAA SUOMI NPP (hereafter SNPP) and NOAA-20, which were launched in October 2011 and November 2017, respectively. Compared to OMI,

OMPS-SNPP is in a relatively lower nadir spatial resolution (OMI: 13×24 km², OMPS-SNPP: 50×50 km²) (de Graaf et al., 2016; Levelt et al., 2006) but has an improved signal-to-noise ratio (González Abad et al., 2016). OMI and OMPS HCHO products share a similar concept and retrieval approach, so the joint evaluation by the two satellites can examine the consistency between OMI and OMPS and, more importantly, provide capability to study HCHO interannual variability on a decadal timescale. Here we use monthly mean HCHO VCD from OMI HCHO VCD retrieval (OMHCHO Version-4) product (González Abad et al., 2022) during 2005-2019 summertime, and OMPS-SNPP Level 2 HCHO total column V1 product (Nowlan et al., 2023) during 2012-2019 summertime, provided by the Smithsonian Astrophysical Observatory.

The OMI and OMPS HCHO retrievals use a three-step procedure to calculate the HCHO VCD (Nowlan et al., 2023). First, the slant column density (SCD) is determined through spectral fitting of a backscattered radiance spectrum collected in the wavelength region of 328.5 to 356.5 nm. This fit uses a daily reference spectrum (one for each cross-track position) determined from radiances collected over a relatively clean area of the Pacific between latitudes 30°S and 30°N. The area used for this reference calculation is referred to as the reference sector. Second, scene-by-scene radiative transfer calculations are performed to determine vertically-resolved scattering weights, which can be used to determine the air mass factor (AMF) in combination with the trace gas profile (Palmer et al., 2001). This AMF describes the path of light and is used for converting the SCD to a VCD (VCD=SCD/AMF). Third, the background reference slant column (SCD_R) in the radiance sector region is determined using a model, to correct the retrieved SCD which is in fact the differential SCD determined from the ratio of the observed radiance and the reference

radiance. A further bias correction (SCD_B) is applied to reduce high-latitude biases, which mostly affect OMPS-SNPP (Nowlan et al., 2023).

To compare with modelled results, OMI and OMPS-SNPP HCHO retrievals are reprocessed following a three-step procedure. This is primarily done to replace the climatology used in the OMI and OMPS-SNPP products with our own GEOS-Chem simulations. First, we remove the data points falling in the following criteria: (1) main quality flag > 0, (2) cloud cover fraction ≥ 40%, (3) solar zenith angle (SZA) ≥ 70°, and (4) Ice/snow flag = 1. After filtering, we regrid the level 2 swath data in the local time window 12:00–15:00 LT to 0.5°×0.625° horizontal resolution. Second, we calculate the air mass factor (AMF_{GC}) based on local GEOS-Chem HCHO vertical profile and satellite scattering weight (Palmer et al., 2001). Third, we calculate the slant column density of HCHO in the reference sector (SCD_{R,SAT}), using modelled HCHO reference sector column and satellite air mass factor over the same location (VCD_{R,GC} and AMF_{R,SAT}) (De Smedt et al., 2018; Zhu et al., 2016):

$$SCD_{R,SAT} = VCD_{R,GC} \times AMF_{R,SAT}$$
 (1)

VCD_{R,GC} is calculated by global monthly climatology of hourly HCHO profiles at the time of overpass, from a 2018 GEOS-Chem high-performance (GCHP) run at $0.5^{\circ}\times0.5^{\circ}$ resolution (Bindle et al., 2021; Eastham et al., 2018). AMF_{R,SAT} is the AMF from the satellite product, which is calculated using the VLIDORT radiative transfer model as described in Nowlan et al. (2023). We rearrange the satellite vertical column as following:

unez retri 160 mol

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Here ΔSCD_{SAT} is the fitted HCHO slant column, SCD_{B,SAT} is the bias correction term for unexplained background patterns in the HCHO retrievals which may be due to instrument or retrieval issues (Nowlan et al., 2023). The single-scene precision of the retrieval is 1×10¹⁶ molecules cm⁻² (absolute) for OMI and 3.5×10¹⁵ molecules cm⁻² for OMPS-SNPP from spectral fitting and 45–105% (relative) from the AMF (González Abad et al., 2015; Nowlan et al., 2023). The spectral fitting error is primarily random in individual measurements, while the AMF error has both random and systematic components. The precision can be improved by spatial and temporal averaging (De Smedt et al., 2008; Zhu et al., 2016; Boeke et al., 2011). Our analyses in this work are based on monthly data, so the absolute uncertainty in HCHO column is reduced to <1×10¹⁵ molecules cm⁻² (De Smedt et al., 2018).

We utilize high-resolution SIF estimates derived from OCO-2 and MODIS (doi: https://doi.org/10.3334/ORNLDAAC/1863, last accessed: August 10, 2022). These datasets provided globally contiguous daily SIF estimates at a spatial resolution of approximately $0.05^{\circ} \times 0.05^{\circ}$ (around 5 km at the equator) and a temporal resolution of 16 days, from September 2014 to July 2020. The dataset was estimated by using an Artificial Neural Network (ANN) trained on the native OCO-2 SIF observations and MODIS BRDF-corrected seven-band surface reflectance along orbits of OCO-2. The ANN model was subsequently used to predict daily mean SIF (mW m⁻²nm⁻¹sr⁻¹) in the gap regions based on MODIS reflectance and land cover. In our study, the OCO-2 SIF estimates are monthly averaged and regridded to $0.1^{\circ} \times 0.1^{\circ}$ spatial resolution for the

comparison with OMI HCHO VCD, and regridded to $2^{\circ} \times 2.5^{\circ}$ spatial resolution when comparing with GEOS-Chem results.

2.2. Global GEOS-Chem simulations

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GEOS-Chem, a 3-D global chemical transport model, is used in this study to examine the spatiotemporal variability of HCHO and VOCs across northern high latitudes. The model is driven by the Modern-Era Retrospective analysis for Research and Applications, Version 2 (MERRA-2), provided by the Global Modeling and Assimilation Office (GMAO) at NASA's Goddard Space Flight Center (Rienecker et al., 2011). GEOS-Chem version 12.7.2 is deployed (doi: 10.5281/zenodo.3701669, last access: August 10, 2022) with an update on cloud chemistry (https://github.com/geoschem/geos-chem/issues/906, last access: August 10, 2022). The simulations encompass 15 summers (1 May to 31 August) from 2005 to 2019, at a horizontal resolution of 2°×2.5° and 72 vertical layers from the surface to 0.01 hPa. For all model runs, we use a standard restart file from the GEOS-Chem 1-year benchmark simulation, followed by an additional spinup period of several days to allow adequate representation of HCHO production and loss in the model.

Biomass burning emissions in our simulation are derived from the Global Fire Emission Database (GFED4.1s) inventory (van der Werf et al., 2017; Randerson et al., 2017). Year-specific GFED4.1s inventory is used in each year of the simulation to make sure the representation of the interannual variability in wildfire emissions. Emissions on a 3-hour basis are obtained from MODIS satellite observations, which provide information on fire detection and burning area (Mu et al., 2011; van

der Werf et al., 2017). The GFED4.1s inventory reports the HCHO emission factor of 1.86 g/kg and 2.09 g/kg dry matter for boreal forest and temperate forest fires (Akagi et al., 2011).

BVOC emissions in the study are calculated online (Emission factor maps computed online) using the Model of Emissions of Gases and Aerosols from Nature (MEGAN, v2.1) (Guenther et al., 2006, 2012) as implemented by Hu et al (2015). Terrestrial vegetation for BVOC emissions is based on the plant functional type (PFT) distribution derived from Community Land Model (CLM4) (Lawrence et al., 2011; Oleson et al., 2013). Utilizing online MEGAN simplifies the investigation of the relationship between BVOC emission patterns and PFTs. CLM4 output (Figure S1) suggests two major PFTs over northern high latitudes: broadleaf deciduous boreal shrubs (mainly over the northern and south Alaska, northern Canada and northern Siberia) and needle leaf evergreen boreal trees (mainly over interior Alaska, northern Canada, south Siberia and the northern part of eastern Europe), both with high emission factors in isoprene and low emission factors in monoterpenes. The southern part of Eastern Europe is dominated by croplands and broadleaf deciduous temperate trees. In this work, 'monoterpenes' from model calculation are lumped monoterpenes, including α -pinene, β -pinene, sabinene and carene.

We conducted a model sensitivity test to assess the difference in BVOC emissions and HCHO dVCD_{Bio,GC} due to online versus offline MEGAN applications. The results of the tests show that the use of online MEGAN has a modest impact on monthly ISOPe and MONOe (25-53% for ISOPe in Alaska, Northern Canada and Eastern Europe, 53% for ISOPe in Siberia; 17-24% for MONOe across the four domains), and provide a similar, isoprene-dominated BVOC emission regime over Alaska, Central Siberia, Northern Canada and Eastern Europe, comparing to results

from using offline MEGAN. The difference in dVCD_{Bio,GC} between using online and offline MEGAN is approximately 13-26%, suggesting a minor impact on dVCD_{Bio,GC} and VCD_{GC} variability over northern high latitudes when using online or offline MEGAN.

In this study, we use the detailed O₃-NO_x-HO_x-VOC chemistry ("tropchem" mechanism) (Park et al., 2004; Mao et al., 2010, 2013), incorporating updates on isoprene chemistry (Fisher et al., 2016). The performance of this version of isoprene chemistry in GEOS-Chem has been extensively evaluated using recent field campaigns and satellite observations over the southeast US (Fisher et al., 2016; Travis et al., 2016), including HCHO production from isoprene oxidation (Zhu et al., 2016, 2020; Kaiser et al., 2018). The ability of GEOS-Chem with this chemistry to reproduce the vertical profiles of HCHO observed during the Alaska summer, as shown in the ATom-1 in-situ campaign, has been demonstrated (Zhao et al., 2022). Under high-NO_x conditions (1 ppbv), HCHO production is rapid, reaching 70-80% of its maximum yield within a few hours, whereas under low-NO_x conditions (0.1 ppbv or lower), it takes several days to reach the maximum yield, and the cumulative yield is approximately 2-3 times lower than that under high-NO_x conditions (Marais et al., 2012).

To examine the influence of different sources on HCHO columns in northern high latitudes, we conducted four GEOS-Chem simulations, as described in Table 1, to separate modelled HCHO total column (VCD_{GC}) into three parts, including the background column (VCD_{0,GC}), biogenic emission induced column (dVCD_{Bio,GC}) as well as wildfire emission induced column (dVCD_{Fire,GC}):

$$VCD_{GC} = VCD_{0,GC} + dVCD_{Bio,GC} + dVCD_{Fire,GC}$$
 (3)

VCD_{0,GC} is the VCD_{GC} from the GEOS-Chem simulation in which both biogenic and wildfire emissions are turned off. VCD_{0,GC}, dVCD_{Fire,GC} and dVCD_{Bio,GC} are derived by Eq. (4a) to (4c):

$$VCD_{0,GC} = VCD_{GC}(BG)$$
 (4a)

$$dVCD_{Fire,GC} = VCD_{GC}(All) - VCD_{GC}(NF)$$
 (4b)

$$dVCD_{Bio,GC} = VCD_{GC}(NF) - VCD_{0,GC}$$
 (4c)

$$dVCD_{Fire,GC}^* = VCD_{GC}(NB) - VCD_{GC}(BG)$$
 (4d)

$$dVCD_{Bio,GC} *= VCD_{GC}(All) - VCD_{0,GC}(NB)$$
 (4e)

To assess the linearity assumption in Eq. (3), we conducted model sensitivity tests over a one-month period to evaluate the disparity between VCD_{GC} and VCD_{0,GC} + dVCD_{Fire,GC}*+ dVCD_{Bio,GC}* (derived from Eq.(4a), (4d) and (4e)). The difference between these two terms is less than 14% in northern high latitudes, suggesting a minor importance of the non-linear effect in this area.

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Figure 1a defines the four domains focused on in this work. The selection of Alaska domain follows Zhao et al (2022); Eastern Europe and Siberia domains follow Bauwens et al (2016); northern Canada domain follows the North America domain in Bauwens et al (2016) but excluded Alaska.

To emphasize the key drivers of HCHO interannual variability, we categorize the years spanning from 2005 to 2019 into two distinct groups: "high HCHO years" and "low HCHO years" within each of the four specified domains. For each domain, the years that have above-average May-August sum of regional-averaged monthly OMI VCD_{SAT,reprocessed} is categorized as "high HCHO years"; those years have the value below average is categorized as "low HCHO years"(shown in Table S1).

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Table 1. Configuration of GEOS-Chem global simulations in this study

Simulations	Biogenic emission	Wildfire
Biogenic + wildfire + Background (All)	On	On
Background (BG)	Off	Off
Biogenic + Background (NF)	On	Off
Wildfire + Background (NB)	Off	On

We use the coefficient of variation (CV) to quantify the interannual variability of summertime HCHO VCD. CV is defined as the ratio of the standard deviation to the mean (CV = $\frac{\sigma}{\mu}$), which is a measure of interannual variability (Giglio et al., 2013). Assuming VCD_{0,GC}, dVCD_{Bio,GC}, dVCD_{Fire,GC} are three independent components of VCD_{GC}, we have $\sigma_{VCD_{GC}}^2 = \sigma_{VCD_{0,GC}}^2 + \sigma_{dVCD_{Bio,GC}}^2 + \sigma_{dVCD_{Fire,GC}}^2$, so the contribution of each component to the CV of VCD_{GC} can be calculated by:

$$CV contribution_{VCD_{0,GC}} = \frac{\sigma_{VCD_{0,GC}}^2}{\sigma_{VCD_{GC}}^2}$$
 (5a)

$$CV contribution_{dVCD_{Bio,GC}} = \frac{\sigma_{dVCD_{Bio,GC}}^2}{\sigma_{VCD_{GC}}^2} \quad (5b)$$

$$CV contribution_{dVCD_{Fire,GC}} = \frac{\sigma_{dVCD_{Fire,GC}}^2}{\sigma_{VCD_{GC}}^2} \quad (5c)$$

3. Evaluation on spatial distribution of HCHO VCD and BVOC emissions

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Figure 1 shows the July mean HCHO VCD over northern high latitudes during 2012-2019, from reprocessed OMI and OMPS-SNPP retrievals, as well as GEOS-Chem model output. We show that OMI and OMPS-SNPP HCHO VCD have consistent spatial patterns and their magnitude agree within 15% (Panel a, b and d). OMPS-SNPP does show lower values in some regions, perhaps due to several cloud and surface reflectance assumptions made in OMPS-SNPP retrievals, or biases that may persist at high-latitudes and large solar zenith angles (Nowlan et al., 2023). While GEOS-Chem well reproduced the spatial pattern of HCHO VCD that OMI and OMPS-SNPP captured (Panel c), we find that GEOS-Chem HCHO VCD is lower than that of OMI by 40%, particularly over wildfire impacted areas (Panel e). The model-satellite discrepancies in wildfire areas can be in part due to model underestimates of VOC emissions and HCHO production from wildfire plumes (Jin et al., 2023), and in part due to the uncertainties in air mass factor calculation for satellite HCHO retrievals in the presence of wildfire smokes (Jung et al., 2019). The model-satellite discrepancies outside wildfire areas could be also due to model underestimates of oxygenated VOCs (OVOCs), biogenic VOC emissions and biases in satellite HCHO retrieval products. For example, Selimovic et al (2022) found that GEOS-Chem underestimates OVOCs, including HCHO, by a factor of 3-12 at Toolik Field Station in Northern Alaska. Stavrakou et al (2015) show model underestimations of biogenic isoprene emission and wildfire emissions over Eastern Europe and Alaska. Recent studies suggest that TROPOMI HCHO retrieval may have a positive bias under low HCHO conditions (Vigouroux et al., 2020). OMPS-SNPP HCHO shows a similar positive bias at clean sites, but has a closer agreement with FTIR HCHO columns at polluted sites (Nowlan et al., 2023; Kwon et al., 2023).

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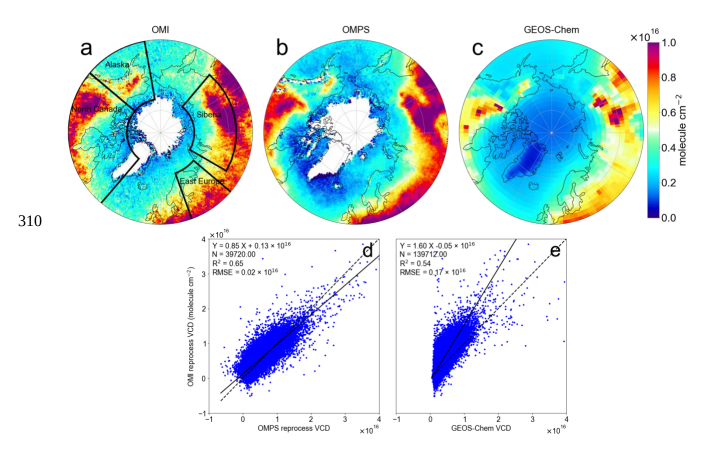


Figure 1. HCHO VCD from OMI, OMPS-SNPP and GEOS-Chem, as well as their linear correlation. (a), (b) and (c) shows Spatial pattern of July mean HCHO VCD from reprocessed OMI, reprocessed OMPS-SNPP and GEOS-Chem over northern high latitudes, in 2012-2019 summers. The black boxes in (a) show the four study domains: Alaska ([50,75] N, [-170, -130] E), Siberia ([57,75] N, [60,140] E), Northern Canada ([50,75] N, [-130, -40] E), Eastern

Europe ([50,71] N, [20,50] E). (d) Scatter plot of monthly HCHO VCD from reprocessed OMI versus reprocessed OMPS-SNPP over continental northern high latitudes in 2012-2019 summers.

OMI and OMPS-SNPP data are regridded to 2° × 2.5° horizontal resolution to matchup with GEOS-Chem pixels. (e) is similar to (d) but shows OMI versus GEOS-Chem.

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We use model sensitivity tests to characterize the spatial variability of HCHO VCD_{GC,0}, dVCD_{Bio,GC} and dVCD_{Fire,GC} over northern high latitudes. Figure 2 shows a similar spatial pattern of VCD_{0,GC} and dVCD_{Bio,GC}, with a distinctive spatial pattern of dVCD_{Fire,GC}. The enhancement of VCD_{0.GC} is mainly shown over Eastern Europe, Eastern Siberia and Central Canada, around 2-4×10¹⁵ molecules cm⁻². dVCD_{Fire,GC} exhibits increases mainly over Alaska, Northern Canada and Central Siberia, with values larger than 5×10¹⁵ molecules cm⁻² at fire hot spots. dVCD_{Bio,GC} spatial pattern corresponds mainly to isoprene emissions over vegetated area, enhances over Eastern Europe (2.4×10¹⁵ molecules cm⁻²) and Eastern Siberia (1.1×10¹⁵ molecules cm⁻²). Model suggests that Eastern Europe is covered by needle leaf evergreen temperate trees and broadleaf deciduous boreal trees, while Eastern Siberia is mainly covered by needle leaf evergreen boreal trees (Figure S1). We note that ΔdVCD_{Bio,GC}:ΔISOPe (Isoprene emission flux. Unit: 10¹⁶ molecules cm⁻² per 1013 atmosC cm-2 s-1) over northern high latitudes is around 0.24, a factor of 10 lower than ΔVCD_{GC}:ΔISOPe over Southeast US (Millet et al., 2008). This indicates a much lower HCHO production efficiency from isoprene oxidation in northern high latitudes compared to mid-latitude, possibly resulting from the availability of NOx, the difference of temperature, photolysis and oxidants level (Marais et al., 2012; Mao et al., 2013; Li et al., 2016; Wolfe et al., 2016).

Our modelled ISOPe is ~1-2 times higher than MONOe (monoterpenes emission) in Alaska, Europe, Northern Canada and central Siberia boreal forest zone, as shown in Figure 4d and 4e. Our model shows comparable isoprene surface mixing ratios with the in-situ measurements along Trans-Siberian Railway within Russian boreal forests (generally <1ppb in our model, and around 0.31–0.48 ppb in the in-situ campaign in Timkovskys et al (2010), both can reach ~4 ppb in Eastern Siberia). Our model also shows comparable monoterpenes surface mixing ratios over Alaskan North Slope (0.009 ppbv in our model and ~0.014 ppbv in Selimovic et al (2022)). Comparing to Stavrakou et al (2018), our modeled ISOPe over Eastern Europe, Alaska and Northern Canada agrees within 20%, but our modeled MONOe is around 40% lower, likely because we are using online MEGAN, different PFT maps and canopy models (Guenther et al., 2012).

A remarkable feature is the heterogeneity of BVOC emissions in northern high latitudes revealed by measurements. We show in Table 2 that while isoprene dominates BVOC emission over the Arctic tundra and broadleaf forests, monoterpene becomes the dominated species over coniferous forests. This includes a large portion over European boreal zone, such as at Hyytiälä in Finland (Rinne et al., 2000; Bäck et al., 2012; Rantala et al., 2015; Zhou et al., 2017; Ciarelli et al., 2024), Bílý Kříž in Czech Republic (Juráň et al., 2017) and Norunda research station in Sweden (Wang et al., 2017). However, this large-scale heterogeneity is not being reproduced by our model. We find from Figure 2f that modeled BVOC emissions are dominated by isoprene in most part of northern high latitudes, except Eastern Siberia and East Greenland. As shown in Figure S1, the isoprene-dominated region is mainly due to broad-leaf deciduous boreal shrubs and needle-leaf evergreen boreal trees that are assumed in the model and exhibits higher isoprene emission factors

than monoterpenes; in contrast, Eastern Siberia is covered predominantly by needle-leaf deciduous boreal trees, leading to higher monoterpenes than isoprene emission (Guenther et al., 2012).

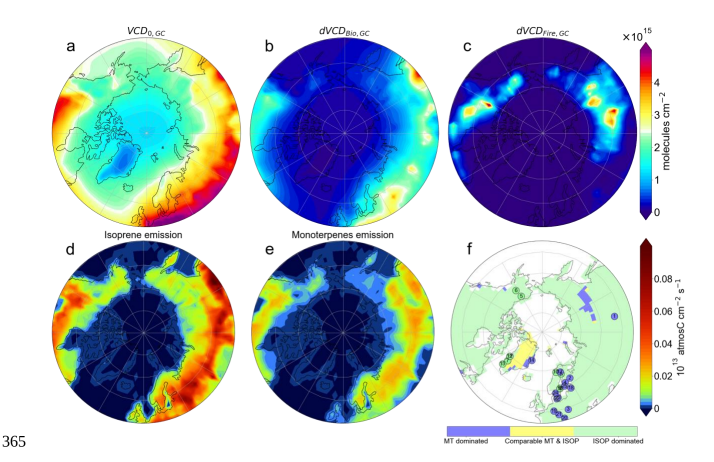


Figure 2. (a) to (e) GEOS-Chem HCHO VCD_{0,GC}, dVCD_{Bio,GC}, dVCD_{Fire,GC}, isoprene and monoterpenes emission fluxes over northern high latitudes, averaged for July from 2005 to 2019. (f) BVOC emission regimes over northern high latitudes, in GEOS-Chem simulation for 2005-2019 summers and from in-situ measurements (references listed in Table 2). Isoprene-dominates regime at a pixel means isoprene emission is significantly higher than monoterpenes emission (p<0.05 in t-test) for May-August in 2005-2019.

Table 2. In-situ measurements of BVOC in Figure 2f

Site name	Lat(°N),	Period of the	Major	Predominant	References
	Lon(°E)	measurement	vegetation	BVOC	
			type at		
			location of		
			measurement		
[1] Irkutsk	53.00,	July 21 to	Boreal	Monoterpenes*	Timkovsky
	102.27	August 4 2008	coniferous		et al., 2010
			forest		
[2]	62.72,	May to October	mixed forest	Monoterpenes*	Hakola et
Pötsönvaar	30.96	1997-1998			al., 2000
a					
[3] Bílý	49.50,	Summer 2009-	Norway spruce	Monoterpenes	Juráň et al.,
Kříž	18.54	2014	forest		2017
[4]	61.84,	1) August 2001	Boreal	Monoterpenes	Spirig et
Hyytiälä	24.29	2) May, 2010 to	coniferous		al., 2004,
		December 2013	forest (Scots		Rantala et
		3) April to	pine (Pinus		al., 2015
		November 2008	sylvestris) and		(emission),
		4) April 2000 to	Norway spruce		Aaltonen et
		April 2002	(Picea abies))		al., 2011
					(emission),

Hakola	et
al., 2003	,

[5] TFS	68.63, -	May to June	Arctic Tundra	Isoprene	Angot et
	149.59	2019			al., 2020;
					Selimovic
					et al., 2022
[6]	64.84, -	August 2016	Needle-leaf	Isoprene*	Zhao et al.,
Fairbanks	147.72		evergreen		2022
			boreal forest		
[7]Kangerl	67.01, -	late June to early	Salix spp.	Isoprene	Vedel-
ussuaq	50.73	August 2013			Petersen et
					al., 2015
[8]	66.61,	May to August	Sub-Arctic fen	Isoprene,	Hellén et
Lompolojä	24.06	2018		Monoterpenes*	al., 2020
nkkä					
[9]	60.08,	May to	mixed forest	Isoprene	Janson et
NOPEX	17.50	September 1995			al., 1999
site					
[10] Abisko	68.35,	June to August	Subarctic wet	Isoprene	Tiiva et al.,
Scientific	18.82	2006-	heath		2008;
		2007,2010-2012			Faubert et

Research			al., 2010;
Station			Valolahti et
			al., 2015
[11] Disko	69.24,	- June to August Subarctic heath Mon	oterpenes (Lindwall
	53.53	2013-2014	et al.,
			2016a)
[12] Disko	69.24,	- June to August Arctic fen Isop	rene (Lindwall
	53.53	2014-2015	et al.,
			2016b)
[13] Nuuk	64.12,	- June to August Subarctic heath Isop	rene Kramshøj
	51.35	2013	et al., 2016
[14]	74.50,	- August 2009 Mesic to dry Mon	oterpenes Schollert
Zackenber	20.50	mixed heath	et al., 2014
g			
[15]	60.08,	June to Dominated by Mon	oterpenes Wang et al.,
Norunda	17.48	September 2013 80-123-years	2017
		old Norway	
		spruce and	
		Scots pine	
[16]	60.08,	June to July 2014 Dominated by Isop	rene (van
Norunda	17.48	80-123-years	Meeningen

			spruce and		
			Scots pine		
[17] Pallas	67.59,	April to August,	Norway Spruce	Sesquiterpenes	Hakola et
Kenttärova	24.15	2020		, Monoterpenes	al., 2023
[18]	58.27,	26 September to	Norway spruce	Monoterpenes	(Bourtsouk
Järvselja,	27.27	17 October 2012			idis et al.,
Estonia					2014b)
[19] Taunus	50.22,	8 April to 11	Norway Spruce	Monoterpenes	(Bourtsouk
Observator	8.43	November 2011			idis et al.,
y, Germany					2014a)
[20]	46.07,	April to May	Norway spruce	Monoterpenes	(van
Ljubljana,	14.50	2016			Meeningen
Slovenia					et al., 2017)
[21]	48.30,	June 2014, 2016	Norway spruce	Monoterpenes	(van
Grafrath,	11.28				Meeningen
Germany					et al., 2017)
[22]	55.67,	July to August	Norway spruce	Monoterpenes	(van
Taastrup,	12.30	2013, July 2014-			Meeningen
Denmark		2016			et al., 2017)

[23]	56.10,	July 2016	Norway spruce	Monoterpenes	(van
Hyltemossa	13.42		with a small		Meeningen
, Sweden			fraction of		et al., 2017)
			Downy birch		
			and Scots pine		
[24]	58.38,	October 2015	coniferous	Monoterpenes	(van
Skogaryd,	12.15		trees,		Meeningen
Sweden			dominated by		et al., 2017)
			Norway spruce		
			and Scots Pine		
[25]	60.38,	July to August,	Norway spruce	Monoterpenes	(van
Piikkiö,	22.50	2014			Meeningen
Finland					et al., 2017)

For the column 'Predominant BVOC', with * means 'Most abundant BVOC in mixing ratio', without * means 'Most emitted BVOC'.

4. Examination on the interannual variabilities of HCHO VCD

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Figure 3 shows that in Alaska, Northern Canada and Siberia, high HCHO years are often associated with strong wildfire VOC emissions (R²=0.78-0.89) and to a lesser extent associated with biogenic VOC emissions (R²=0.21-0.47). The interannual variability of wildfire VOC emission is further supported by CO emissions from both GFED4 and satellite-based estimation (Yurganov and Rakitin, 2022). The high correlation between OMI HCHO VCD and GFED wildfire VOC

emissions in Alaska, Siberia and Northern Canada indicates a strong wildfire impact on interannual variabilities of HCHO VCD in these domains. In Eastern Europe, high HCHO years are associated with large biogenic emissions (With wildfire VOC emissions: R²=0.51; With biogenic emissions: R²=0.72), indicating the important role of biogenic emission in interannual variability of HCHO in Eastern Europe.

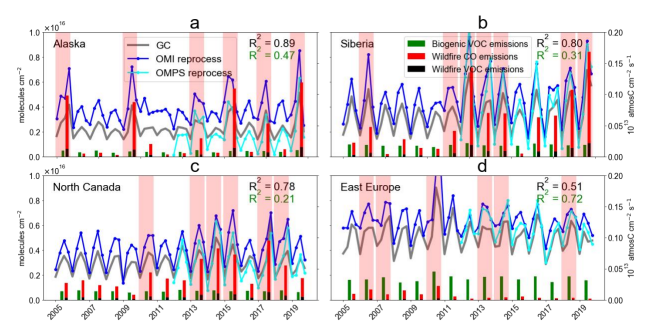


Figure 3. Timeseries of HCHO VCD, biogenic and wildfire emissions over (a) Alaska, (b) Siberia, (c) Northern Canada and (d) Eastern Europe, May 1-August 31, 2005-2019. The blue lines are monthly HCHO VCD from reprocessed OMI, cyan lines are from reprocessed OMPS-SNPP, grey lines are from GEOS-Chem. Red and black bars are area-normalized wildfire CO and VOC emissions during the summer of each year; green bars are area-normalized biogenic VOC emissions. Wildfire emissions are calculated from GFED4.1s inventory; biogenic VOC emissions are calculated by MEGAN2.1 model. Pink shade indicates high HCHO VCD years (definition see Sect. 2.2 and Table S1). The R² between reprocessed OMI HCHO VCD and biogenic VOC emission (green) / wildfire VOC emission (black) is shown at top right of each panel.

Figure 4a to 4c shows that wildfire is the main driver of HCHO VCD_{GC} interannual variability over Siberia, Northern Canada and Alaska. In low HCHO years of these three domains, dVCD_{Fire,GC} contribution ~2-11% of HCHO total column, less than VCD_{0,GC} and dVCD_{Bio,GC}; in high HCHO years, dVCD_{Fire,GC} contribution to total column rises to ~20-34%. This is consistent with Figure 3 that HCHO VCD interannual variability have significantly higher correlations with wildfire emissions (R²=0.78-0.89) than with biogenic emission (R²=0.21-0.47) over Siberia, Northern Canada and Alaska. These findings highlight the role of wildfire in driving HCHO interannual variability in the three domains.

In Eastern Europe, biogenic emission and background HCHO accounts for the majority of HCHO VCD interannual variability, largely due to the relatively higher surface temperature, stronger photolysis, higher oxidants level and higher availability of NO_x than the other three domains. In regional scale, BVOC emissions and methane oxidation with hydroxyl radicals (OH) both depend on temperature (Guenther et al., 2012; Holmes et al., 2013). In Figure 4d, the surface temperature in Eastern Europe is higher than that in Alaska, Northern Canada and Siberia by 5-7 K, leading to an increase in BVOC emissions and VCD_{0,GC} through methane oxidation. HCHO VCD is further enhanced through the higher NO_x level (0.4-1ppbv) in Eastern Europe than in other three domains (0.1-0.5ppbv), as HCHO yield from isoprene photooxidation increases with NO_x level. The high NO_x level in Eastern Europe results from its large urban areas and high anthropogenic emissions. The large contribution of BVOC to HCHO VCD is consistent with Figure 5, which shows the CV of dVCD_{Bio,GC}+VCD_{0,GC} accounts >90% of VCD_{GC}'s CV in Eastern Europe. Similarly, Figure 3d shows that biogenic emission has a higher correlation (R² = 0.72) with VCD_{GC} than wildfire

emission does (R²=0.51). These results suggest that biogenic emission and background are the main contributors of HCHO interannual variability in Eastern Europe.

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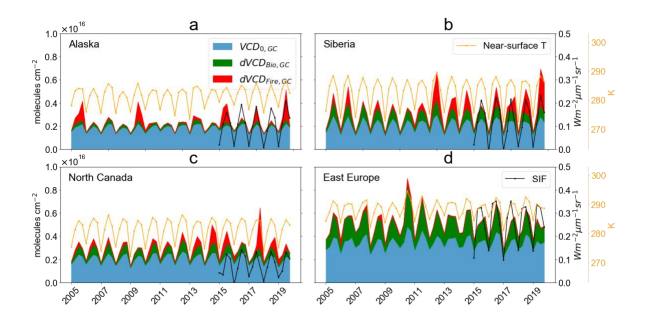


Figure 4. Interannual variability of monthly HCHO VCD_{0,GC}, dVCD_{Bio,GC} and dVCD_{Fire,GC} as well as near-surface temperature over (a) Alaska, (b) Siberia, (c) Northern Canada and (d) Eastern Europe, in 2005-2019 summers. In each year, only the monthly values in May, June, July and August are shown. The indigo, green and red shades are background HCHO VCD_{0,GC}, dVCD_{Bio,GC} and dVCD_{Fire,GC}, based on GEOS-Chem sensitivity tests (Table 1). The orange curves are monthly surface temperature from MERRA-2 dataset. The black curves are OCO-2 monthly SIF.

We further examine the contribution from background, biogenic and pyrogenic emissions to the interannual variability of HCHO VCD_{GC} over each region. We find from model results that biogenic emission and background signal contributes to 90% of the interannual variability of HCHO VCD_{GC} in Eastern Europe, while wildfire accounts for over 90% of CV in Alaska, Siberia

and Northern Canada, consistent with previous work (Stavrakou et al., 2018; Zhao et al., 2022).

We use Mann-Kendall test, a non-parametric statistical test used to detect trends in time series data, to test the significance of the trend of monthly HCHO VCD_{GC} time series over a specific domain (Gilbert, 1987). We found no significant trend of HCHO VCD_{GC} over Eastern Europe, Northern Canada and Alaska from either satellites or model. On the other hand, we find the trend of HCHO VCD_{GC} over Siberia is significant (p<0.05) and increasing (1.7% per year). VCD_{0,GC} and dVCD_{Bio,GC} show no significant trend, while the trend of dVCD_{Fire,GC} is significant and increasing in Siberia (12% per year), suggesting that wildfires are responsible for the VCD_{GC} trends in Siberia. In contrast to Bauwens et al (2016), We find that HCHO VCD_{GC} trend over Siberia is largely driven by the increasing wildfires in recent years, and to a lesser extent by biogenic VOC emissions, highlighting the important role of wildfires on HCHO VCD interannual variability.

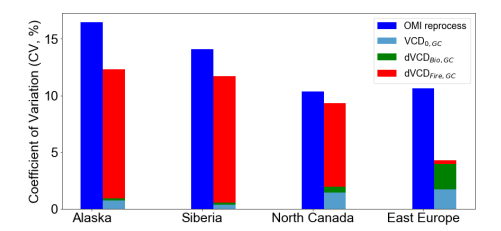


Figure 5. Coefficient of Variation (CV) of OMI HCHO VCD, modelled HCHO VCD_{0,GC}, $dVCD_{Bio,GC}$ and $dVCD_{Fire,GC}$ in summertime of 2005-2019.

5. SIF evaluation on dVCD_{Bio,GC} interannual variability

In Figure 6a to 6d, we find a good linear relationship (R=0.6-0.7) between OCO-2 monthly SIF and dVCD_{Bio,GC} at Alaska, Siberia, Northern Canada and Eastern Europe. Foster et al (2014) show a high linear correlation between seasonal variation of satellite HCHO column (fire free) and GPP in northern high latitudes. This is consistent with our finding over most continental areas in northern high latitudes (Figure S2), since SIF is a widely used proxy of GPP (Frankenberg et al., 2011). In Figure 6g to 6j, SIF and ISOPe show a linear relationship when SIF is within 0-0.25 Wm⁻² μ m⁻¹sr⁻¹ but tend to decouple when SIF > 0.25 Wm⁻² μ m⁻¹sr⁻¹, possibly due to the different temperature optimums of isoprene emission and photosynthesis (Harrison et al., 2013; Zheng et al., 2015).

Despite the difference in distribution of vegetation types, the dVCD_{Bio,GC}-SIF slope is homogeneous over Siberia, Northern Canada and Eastern Europe (slope=0.28-0.45, unit:10¹⁶ molecules cm⁻² per Wm⁻²μm⁻¹sr⁻¹), suggesting SIF as a tool to understand biogenic HCHO variability in these regions. The dVCD_{Bio,GC}-SIF slope in Alaska is 3-5 times lower than other three domains, which warrants further investigation. In contrast to high latitudes, we find that both ISOPe:SIF slope and dVCD_{Bio,GC}:SIF slope are a factor of 2-10 times higher in Southeast US and Amazon (Figure 6e-6f, 6k-6l) than in northern high latitudes, indicating that the dVCD_{Bio,GC}-SIF slope over northern high latitudes and lower latitudes could be very different.

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SIF offers an independent evaluation on the interannual variability of HCHO dVCD_{Bio,GC}. As SIF showing a linear relationship with dVCD_{Bio,GC} in northern high latitudes (Figure 6a to 6d), it is reasonable to infer from Figure 4 that the low interannual variability shown in SIF (CV=1-9%) is

expected for dVCD_{Bio,GC} (CV=1-2%) in Alaska, Siberia and Northern Canada. In contrast, we find that dVCD_{Fire,GC} has a much weaker correlation with SIF (Figure S2c) and shows a higher interannual variability (CV=8-13%). As wildfire emission is highly correlated (R²=78-89%) with OMI HCHO VCD over Northern Canada, Siberia, and Alaska (Figure 3), the high interannual variabilities of OMI HCHO VCD (CV=10-16%) in these domains are likely driven by wildfires instead of biogenic emissions.

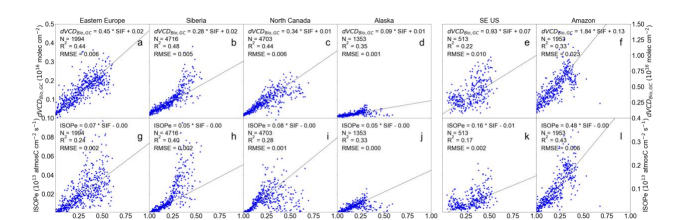


Figure 6. Scatter plot of monthly OCO-2 SIF versus GEOS-Chem HCHO dVCD_{Bio,GC} and isoprene emission fluxes in the four study domains plus Southeast US ([26, 36]°N, [-100,-75]°E) and Amazon([-20,-5]°N, [-75,-40]°E), from May to August in 2015-2019. OCO-2 SIF is regridded to 2°×2.5° spatial resolution. Only continental pixels of SIF-dVCD_{Bio,GC} and SIF-ISOPe matchups are used to plot. Before plotting, data matchups are binned by SIF, using a bin size of 0.001 Wm⁻²μm⁻¹sr⁻¹. Linear regression is shown as the black dash in each panel, calculated for SIF within 0-0.25 Wm⁻²μm⁻¹sr⁻¹. Amount of binned data pairs (N), R-Squared (R²), Root Mean Square Error (RMSE) are calculated based on binned data across all ranges.

6. Conclusions and discussions

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We use reprocessed new retrievals of HCHO from OMI and OMPS-SNPP to evaluate the interannual variability of HCHO VCD from GEOS-Chem over northern high latitudes in 2005-2019 summers. The reprocessed OMI and OMPS-SNPP HCHO VCDs show a high consistency in the spatial pattern and interannual variability. GEOS-Chem reproduced the interannual variability of HCHO VCD but the magnitude is biased low comparing to satellite retrievals.

Our modeled HCHO VCD can be biased low, due to large underestimate of HCHO production and emission factor in wildfire smokes. Previous in-situ campaigns show underestimated emission factors of VOCs in GFED4.1s emission inventory for temperate forests in western US (Liu et al., 2017; Permar et al., 2021), while the bias in VOC emission factor in boreal forest wildfires remains unclear. HCHO underestimation can also be due to the missing HCHO secondary production in wildfire-impacted conditions (Liao et al., 2021; Jin et al., 2023). GEOS-Chem is found to underestimate oxygenated VOCs by a factor of 3 to 12 in some Arctic regions, which could contributes to the bias in modelled HCHO in northern high latitudes (Selimovic et al., 2022). More measurements in Arctic region are needed to reconcile the model-observation discrepancies.

Wildfire accounts for the majority of HCHO interannual variability in Alaska, Northern Canada and Siberia. Compared to biogenic emissions and background HCHO, wildfire emission shows a better correlation with HCHO VCD, despite that biogenic and background HCHO can dominate HCHO VCD in low HCHO years of these three regions. We also find an increasing trend (p < 0.05) in wildfire emission and HCHO VCD over Northern Canada and Siberia. With rapid Arctic warming, wildfire frequency and intensity rises rapidly in recent decades and near future (Descals

et al., 2022). We expect wildfire continues to dominate HCHO interannual variability in the three regions.

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Eastern Europe is the only one of the four studied regions where HCHO interannual variability is dominated by biogenic emission and background HCHO. This is due to a combination of lower wildfire activities, higher surface temperature and anthropogenic NO_x emissions in this region. No significant trend of biogenic emission, biogenic-related HCHO and background HCHO are found in the four regions during summertime of 2005-2019. However, model estimate of HCHO from biogenic emissions are largely uncertain, as model calculated VOC speciation is at odds with field measurements (Figure 2f and Table 2). Previous work shows good performance of model in capturing long-term variability of biogenic emission in response to climate variables (Stavrakou et al., 2018), but model underestimates biogenic and fire emissions over northern high latitudes, especially over Eastern Europe and Alaska (Stavrakou et al., 2015). Future research is warranted to examine the HCHO signal from biogenic emissions in this region.

The OCO-2 satellite SIF provides an additional constraint on the interannual variability of biogenic emissions and is independent of wildfire emissions. As a proxy of vegetation photosynthesis and GPP, SIF is expected to have a good correlation with isoprene emission and HCHO VCD in the northern boreal regions, though this correlation can be worse in mid-latitudes and tropical region (Foster et al., 2014). We show a tight and homogeneous linear relationship (R=0.6-0.7) between SIF and dVCD_{Bio,GC} at northern high latitudes, suggesting that SIF may serve as a tool to understand biogenic HCHO variability in this region.

Code and data availability.

The L2 V1 **OMPS-SNPP HCHO** product is available at 545 https://disc.gsfc.nasa.gov/datasets/OMPS_NPP_NMHCHO_L2_1/summary (González Abad, 2022). The OMI L2 **HCHO** product is available at https://waps.cfa.harvard.edu/sao_atmos/data/omi_hcho/OMI-HCHO-L2/ . The OCO-2 SIF is available at https://daac.ornl.gov/cgi-bin/dsviewer.pl?ds id=1863 (Yu et al., 2021). Data used in this work is available at https://doi.org/10.6084/m9.figshare.23599566.v1 (Zhao, 2023a). Data 550 processing and plotting codes are available at https://doi.org/10.5281/zenodo.8094844 (Zhao, 2023b). The GEOS-Chem model is publicly available at: https://doi.org/10.5281/zenodo.3701669 (GEOS-Chem, 2020).

Supplement.

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555 The supplement related to this article is available online at: https://doi.org/10.6084/m9.figshare.25130813.v1.

Author contributions.

TZ and JM designed the research, performed the simulations and conducted the analysis. ZA, GGA and CN provided OMI and OMPS data. YZ helped process and analyze the data. TZ and JM wrote the paper with all co-authors providing input.

Competing interests.

The contact author has declared that neither they nor their co-authors have any competing interests.

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