



Wintertime photochemistry of acyl peroxy nitrates and ozone in South Korea during the ASIA-AQ campaign

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40 **Abstract**



Wintertime photochemical air pollution in East Asia remains poorly constrained despite its impact on regional air quality. Sources and formation pathways of acyl peroxy nitrates (PANs) and ozone (O₃), key photochemical products, are not well understood, hindering effective mitigation strategies. We investigate PANs and O₃ over South Korea using observations from the ASIA-AQ campaign (February–March 2024). PANs reached 5.5 ppbv, strongly correlating with formaldehyde and particulate matter, indicating active winter photochemistry. Median PANs were higher in the mid-southern peninsula (MS; 987 pptv) and Yellow Sea (1197 pptv) than the Seoul Metropolitan Area (842 pptv). Elevated homologue-to-acetyl peroxy nitrates ratios over the MS, with enhanced acryloyl peroxy nitrates, acrolein, and ethylene oxide, provided tracers for petrochemical emissions and their impacts. Acetaldehyde contributed 53–80% of PAN production. Ethanol was a major precursor of acetaldehyde (~50%). Strong correlations ($r^2 > 0.8$) of ethanol and halocarbons indicate industrial and solvent sources under-represented in inventories. Formaldehyde and C₂+ aldehydes contributed ~30% to ozone production. Low ozone production efficiency (< 10) and radical termination dominated by nitric acid and PANs (> 80%) indicate VOC-limited conditions. The fractional PANs contribution to NO_x loss increased with decreasing OH reactivity ratio of NO₂ to aldehydes, suggesting spatial increases in ozone production following NO_x reductions. These findings demonstrate that a comprehensive understanding of VOC oxidation, particularly oxygenates from industrial sources, is essential for representing winter photochemistry. PANs measurements provide critical constraints on oxidation processes and their implications for emission control.

1 Introduction

Wintertime photochemical smog remains a persistent air quality challenge in East Asia. Despite reduced solar radiation and temperatures, observations indicate active oxidation of volatile organic compounds (VOCs) and nitrogen oxides (NO_x = NO + NO₂) by atmospheric radicals, especially the hydroxy radical (OH) (Tan et al., 2018; Lu et al., 2019; An et al., 2019). The photooxidation of VOCs and NO_x (hereafter VOC–NO_x photochemistry) produces a suite of oxidized VOCs species when sufficient OH levels are sustained by photolysis of nitrous acid (HONO) and ozone (O₃). Products include aldehydes (Apel et al., 2010), as well as NO_x oxidation products (NO_z) such as nitric acid (HNO₃), acyl peroxy nitrates (PANs; RC(O)OONO₂), alkyl nitrates (ANs; RONO₂) and particulate nitrate (pNO₃⁻) (Guo et al., 2014; Tan et al., 2018; Fu et al., 2020; Lee et al., 2021).

Understanding the atmospheric impacts of VOC–NO_x photochemistry requires an accurate representation of emissions and oxidation pathways (Bates et al., 2021; von Schneidmesser et al., 2023). A growing body of observations in East Asia points to high VOC levels influenced by complex emissions from combustion (e.g., vehicle exhaust and biomass burning) and non-combustion sources (e.g., volatile chemical products; VCPs), both of which are poorly constrained in inventories (McDonald et al., 2018; Li et al., 2019; Simpson et al., 2020; Travis et al., 2024). Uncertainties in emissions, and consequently in post-emission atmospheric processing, translate to uncertainties in the simulation of photochemical products, including major smog pollutants such as O₃ and fine particulate matter (PM_{2.5}). Reducing these uncertainties is therefore critical



for predicting the extent of air pollution and formulating mitigation strategies for present environmental issues such as decadal rise in average surface O₃ concentrations, increasing winter–spring O₃ levels, and severe winter haze events (Cooper et al., 2020; Li et al., 2020, 2021; Zhai et al., 2019).

VOC emission reductions have been proposed to mitigate O₃ and particulate pollution (i.e., VOC-limited chemical regime) in winter–spring East Asia, but the complexity in VOC emissions and chemical evolution complicate identification of which VOCs contribute to photochemistry and pollutant formation (Li et al., 2021; Schroeder et al., 2020; Nault et al., 2018). An experimental approach is to examine observed secondary products of VOC oxidation (e.g., Ryerson et al., 2003). Among these, formed via VOC–NO_x photochemistry, PANs are effective tracers of VOC oxidation and often a major component of NO_z (Roberts et al., 2007). Recent field observations across East Asia, including China and South Korea, underscore the diagnostic utility of PANs as tracers of VOC–NO_x photochemistry. For example, in China, wintertime acetyl peroxyxynitrate (PAN), the simplest and most abundant member of PANs, strongly correlates with PM_{2.5}, often more so than with O₃, partly due to titration of O₃ by elevated NO_x (Qiu et al., 2019, 2020; Lu et al., 2019; Xu et al., 2021). Here, PAN serves as a more useful indicator of photochemistry than O₃ and provides observational evidence for the contribution of atmospheric chemistry to particulate pollution. Similarly, in the Seoul Metropolitan Area (SMA), home to 50% of South Korea's population, springtime production of O₃ was found to be dominated by local oxidation of small (<C₅) anthropogenic alkenes and aromatics (>C₇) such as toluene and xylenes from traffic and solvent emissions. Organic aerosol (OA) production was driven by oxidation of aromatics and other VOCs with semi- and intermediate volatility (SVOCs and IVOCs) (Simpson et al., 2020; Schroeder et al., 2020; Nault et al., 2018). The dominant role of secondary formation is further demonstrated by the co-occurrence of OA and O₃ with short-lived photochemical tracers including formaldehyde (CH₂O) and PAN (Schroeder et al., 2020; Nault et al., 2018).

More recently, an evaluation of Korea–United States Air Quality (KORUS-AQ) observations by Nault et al. (2024) showed that approximately half of the species comprising PANs predicted by near-explicit box model simulations were unmeasured despite their importance in O₃ and radical chemistry in the SMA. To reconcile the PANs budget, GEOS-Chem transport model simulations required increased VOC emissions from non-combustion sources, such as VCPs, and revised chemical mechanisms (Bey et al., 2001; Travis et al., 2024). Consistent with this, in New York City (NYC), PAN formation in WRF-Chem simulations was highly sensitive to oxygenated VCPs such as ethanol and isopropanol whose inclusion increased predicted PAN by 15–20% (Grell et al., 2005; Coggon et al., 2021). These studies demonstrated difficulties in constraining both the magnitude and speciation of PANs, and their utility as VOC–NO_x photochemical tracers in polluted urban environments with complex VOCs.

A large data set of PANs observations has been accumulated globally, spanning regions from remote hemispheric backgrounds to heavy pollution (Roberts et al., 2007; Fischer et al., 2014; Lee et al., 2025). However, such observations in East Asia, including South Korea, remain scarce. To address this gap, *in situ* airborne observations onboard the NASA DC-8 were conducted over South Korea in February and March 2024 as part of the Airborne Satellite Investigation of Asian Air Quality (ASIA-AQ) campaign. A key objective was to assess South Korean air quality in winter and build upon KORUS-AQ



findings (Crawford et al., 2021). The ASIA-AQ instrument suite includes measurements of PANs, their oxygenated VOC (OVOC) precursors (e.g., aldehydes, alcohols, and esters), and aromatic oxidation products. To our knowledge, no prior studies have provided such a comprehensive measurement of those compounds. Therefore, the ASIA-AQ observations offer a unique opportunity to characterize wintertime photochemical processing in East Asia.

This work investigates wintertime photochemistry in South Korea, focusing on the chemistry and distribution of PANs. We report measurements of PANs, including rarely observed homologues such as acryloyl peroxyxynitrate (APAN) and benzoyl peroxyxynitrate (PBzN). Combining kinetic calculations and 0-D box modeling, highly constrained by observations, we examine key factors controlling PAN and O₃ production. For this examination, we focus on diagnosing instantaneous PAN and O₃ production using observed precursors across different regions in South Korea. To systematically represent spatial and pollution-dependent variations in photochemistry, we analyze the fractional contribution of NO_x loss pathways to either PANs or HNO₃ formation as well as ozone production efficiency (OPE) as a function of the probability of OH reacting with NO₂ or aldehydes.

2 Methods and data analysis

2.1 Brief overview of the ASIA-AQ campaign

The ASIA-AQ campaign was conducted over the Philippines, South Korea, Thailand, and Taiwan from 6 February to 27 March 2024. Airborne measurements were primarily obtained from the NASA DC-8 and G-III, complemented by satellite, ground-based, and additional aircraft observations from domestic collaborators (ASIA-AQ, 2023). This work focuses on five DC-8 research flights (RFs) over South Korea and the Yellow Sea from 17 February to 11 March, during cold conditions (median temperature of 4°C at altitudes below 500 m).

During ASIA-AQ, DC-8 flights departed from Osan Air Base and followed consistent transects at altitudes ranging from ~200 m to 2.5 km. The near-constant flight paths were performed to obtain consistent and representative sampling of the South Korean atmosphere. Each RF included 4–6 low approaches at Seoul and Gimpo Airports to capture vertical distributions over urban areas, totaling 27 low approaches over the SMA during the campaign. To assess regional air pollution, the DC-8 flights included sampling over relatively remote regions such as the Yellow Sea and the mid- and southern regions of the peninsula. Measurement locations and conditions during the campaign are shown in Fig. S1 in the Supporting Information (SI).

2.2 TD-CIMS measurements

A comprehensive suite of chemical, optical, and meteorological measurements was obtained aboard the DC-8 aircraft during the ASIA-AQ campaign. Table S1 summarizes the measurements used in this study, including instrumental techniques, uncertainties, and references. Of the measured compounds, this study focuses primarily on PANs measured by the Georgia Tech Thermal Dissociation-Chemical Ionization Mass Spectrometer (GT TD-CIMS) with a Time-of-Flight (ToF) mass



analyzer (ToFWerk/Aerodyne). Building on established methods (Slusher et al., 2004; Zheng et al., 2011; Lee et al., 2020), this deployment incorporated several key modifications, as shown in Fig. S2. A major improvement was the replacement of the quadrupole mass filter with a ToF analyzer (resolving power ~ 5000 m/ Δ m), which enhanced mass resolution and thereby improved the selective measurement of PAN homologues. As in our prior DC-8 deployments, quantification of PANs was further supported by the continuous addition of isotopically labeled PAN ($C^{13}H_3C^{13}(O)OONO_2$) for calibrations and periodic NO addition to react away peroxyacyl radicals in the thermal dissociation (TD) region for background determination. In addition, post-campaign laboratory experiments were carried out to constrain sensitivities of PAN homologues relative to that of PAN, following the method described in Roberts et al. (2022).

An additional modification to the GT TD-CIMS was the use of an automated variable orifice upstream of the heated Teflon inlet (i.e., TD region), replacing the in-line pressure controller used in the KORUS-AQ and Atmospheric Tomography (ATom) campaigns (Fig. S2). The variable orifice, previously applied to regulate flow tube pressure across altitude ranges (Chen et al., 2016), maintained constant pressure in both the TD and Ion Molecule Reaction (IMR) regions during ASIA-AQ. This also reduced interactions with metal surfaces compared to the previous inlet system (Lee et al., 2020). While wall losses for most PANs are minimal, this configuration is expected to improve the measurements of PBzN for which potential loss on Teflon inlet wall has been reported (Zheng et al., 2011; Liu et al., 2019). It should be noted that such potential effects were further resolved by applying homologue-specific sensitivities during data analysis.

Finally, the CIMS instrument utilized a vacuum ultraviolet (VUV) lamp (Heraeus, PKS 106) as an ion source (Ji et al., 2020), eliminating the safety and regulatory constraints associated with polonium-210 sources (1.5–20 mCi) used in previous flight instruments (e.g., Lee et al., 2020, 2022). For a campaign such as ASIA-AQ, with the extended duration and multiple deployment sites, the VUV lamp reduces logistical demand in contrast to radioactive sources that require regulatory oversight (Lee et al., 2020).

2.3 VOCs measurements and data analysis

This study includes kinetic calculations and 0-D box modeling, both highly constrained by observations, including an extensive suite of hydrocarbons and OVOC measurements, on the NASA DC-8. During ASIA-AQ, VOCs were measured by the NSF NCAR Trace Organic Gas Analyzer with a time-of-flight mass spectrometer (TOGA-TOF), the University of California, Irvine Whole Air Sampler (WAS), and the University of Oslo FUSION proton transfer-reaction time-of-flight mass spectrometer (PTR-ToF-MS; Reinecke et al., 2023). Among these, the TOGA-TOF, which utilizes fast *in situ* gas chromatography coupled with a high-resolution time-of-flight electron impact mass spectrometer served as the primary dataset (Apel et al., 2015; Hornbrook et al., 2016). Analyses incorporating VOC observations utilized data merged to the TOGA sampling interval (35 seconds of integrated sampling every two minutes). For analyses using only PANs data (Sect. 3.1), measurements were merged to a 1-second time base. All DC-8 measurement data are publicly available through the NASA data archive (<https://doi.org/10.5067/SUBORBITAL/ASIA-AQ/DATA001>).



170 To assess regional variability in atmospheric composition and photochemistry, the time-averaged data were categorized into three regions based on geographic boundaries: Seoul and Suburban (SMA), Mid and South (MS), and Yellow Sea (YS) (Fig. S3a). In addition, we incorporated 1 km × 1 km gridded data from the Statistical Geographic Information Service (SGIS) of the Korean government (<https://sgis.kostat.go.kr>) to complement chemical observations with spatially resolved information relevant to anthropogenic activities. The statistical data includes industrial facilities, classified by sector
175 (Fig. S3 b–c), as well as population density. For this analysis, industrial sectors included manufacturing, power generation, and utilities/management. The population and industrial facility densities were averaged within a 5 km radius of each DC-8 measurement point, denoted as PD_{5-km} and ID_{5-km}, respectively, and used to support qualitative interpretation of pollutant distribution using chemical tracer measurements.

2.3.1 Kinetic calculations of instantaneous rates and reactivity

180 A series of kinetic calculations were conducted to estimate instantaneous production rates of O₃ and reactive nitrogen (NO_y), including nitric acid (HNO₃) and PANs, based on observed concentrations, photolysis frequencies, and established kinetic parameters. These estimates were integrated with observations and box model simulations to investigate O₃ and NO_z formation during the campaign.

As daytime oxidation of atmospheric species is primarily initiated by reaction with OH, instantaneous production
185 rates, considered here, depend on OH reactivity. In this work, the OH reactivity of an individual compound X_i (R_{X_i}) is defined as the product of its observed concentrations and its corresponding OH rate coefficients (k_{OH}). The summed OH reactivity (R_X) is then obtained by adding the individual contributions (Eq. (1)).

$$R_X = \sum_i R_{X_i}, \text{ where } R_{X_i} = k_{OH+X_i}[X_i] \quad (1)$$

190 The instantaneous production rate of ozone (P(O₃)) can be approximated by the rate at which peroxy radicals (HO₂ and RO₂) react with NO (Eq. (2)) and was used to calculate modeled P(O₃). Given the absence of peroxy radical measurements during ASIA-AQ, we estimate P(O₃) using Eq. (3), following Rosen et al. (2004) and Perring et al. (2013), which represent ozone production as a function of R_{VOCs}, OH concentration, speciated radical yields (γ_i) and alkyl nitrate branching ratios (α_i), as summarized in Table S2. A constant OH concentration of 2 × 10⁶ molecules cm⁻³ was used for the base case to generalize
195 the analysis and enable application across a broader observational dataset than model simulations that require fully synchronized inputs. Modeled OH concentrations were used when comparing kinetic calculations to model simulations.

$$P(O_3) \cong \sum_i k_{RO_{2,i}+NO}[RO_{2,i}][NO] + \sum k_{HO_2+NO}[HO_2][NO] \quad (2)$$

$$P(O_3) \cong \sum_i \gamma_i(1 - \alpha_i)[OH]R_{VOC_i} \quad (3)$$



Similarly, production rates of HNO_3 $P(\text{HNO}_3)$ and PANs $P(\text{PANs})$ are estimated using Eqs. (4)–(5), respectively. In
 200 Eq. (5), β_{PAN} represents the fraction of acyl peroxy radical (PAs) reacting with NO_2 relative to the total loss pathways, and
 α_{HA} is the branching ratio for hydrogen abstraction of the corresponding aldehydes. The OH reactivity from aldehyde
 precursors of PANs (R_{ALD}) is calculated using observed mixing ratios of acetaldehyde, propanal, butanals, acrolein and
 benzaldehyde. The photolytic loss of OVOC, $j_i[\text{OVOC}_i]$, is the product of measured photolysis frequencies and OVOCs
 concentrations (acetone, methyl ethyl ketone, and biacetyl), weighted by the photolytic yield of acetyl peroxy (PA) radical
 205 ($\gamma_{i,\text{PA}}$), generally unity except for biacetyl, which yields two PA radicals. Based on the rate estimates, ozone production
 efficiency (OPE) is approximated as the ratio of $P(\text{O}_3)$ to the sum of HNO_3 and PANs production rates (Eq. (6)).

$$P(\text{HNO}_3) = [\text{OH}]R_{\text{NO}_2} \quad (4)$$

$$P(\text{PAs} + \text{PANs} \cong \text{PANs}) = \sum_i \beta_{\text{PANs}} (\alpha_{\text{Hydrogen Abstraction},i} [\text{OH}]R_{\text{ALD},i} + \gamma_{i,\text{PA}} j_i [\text{OVOC}_i]), \quad (5)$$

$$\text{where } \beta_{\text{PAN}} = \frac{k_{\text{PA}+\text{NO}_2} [\text{NO}_2]}{k_{\text{PA}+\text{NO}_2} [\text{NO}_2] + k_{\text{PA}+\text{NO}} [\text{NO}]}$$

$$\text{OPE} = \frac{P(\text{O}_3)}{P(\text{HNO}_3) + P(\text{PANs})} \quad (6)$$

2.3.2 Steady state 0-D box modeling

We used a zero-dimensional box model (F0AM; The Framework for 0-D Atmospheric Modeling; Wolfe et al., 2016)
 210 to simulate production rates of O_3 and NO_z . The model was constrained using synchronous measurements of trace gases and
 meteorological parameters that were averaged to the TOGA sampling interval (TOGA-merge). A complete list of observational
 constraints is provided in Table S3. To interpret airborne measurements under rapidly changing spatial and temporal
 conditions, we used the flight steady-state method (Crawford et al., 1999). This approach is widely used in airborne studies
 and is particularly suited for estimating concentrations of short-lived species (e.g., radicals) and diagnosing instantaneous
 215 photochemistry (Olson et al., 2006; Schroeder et al., 2020). In this work, each model step is treated as an independent steady-
 state run incorporating the near-explicit chemical mechanism (the Master Chemical Mechanism (MCM) v3.3.1; Jenkin et al.,
 2015) to iterate through full diurnal cycles until unconstrained species (e.g., radicals) converged within 1% (e.g., Nault et al.,
 2024). A similar approach has been previously applied to investigate radical chemistry, ozone production and reactive nitrogen
 processing for KORUS-AQ (Brune et al., 2022; Schroeder et al., 2020; Nault et al., 2024). To account for differences between
 220 observed and modeled j values, we apply a scaling factor to all j values based on the averaged ratio of measured to MCM-
 parameterized j_{NO_2} and j_{O_3} . In addition, a first-order loss is applied to all species, equivalent to a 12-hour lifetime, following
 Brune et al. (2022) and Nault et al. (2024). However, due to the short atmospheric lifetime of the species considered here and
 the use of highly constrained input for instantaneous rate calculations, this loss term has a minimal influence ($< \sim 5\%$ variability
 for a factor of two change in loss rate) on the results in this study (Fig. S4).



225 3 Observations of PANs over the Korean Peninsula and the Yellow Sea

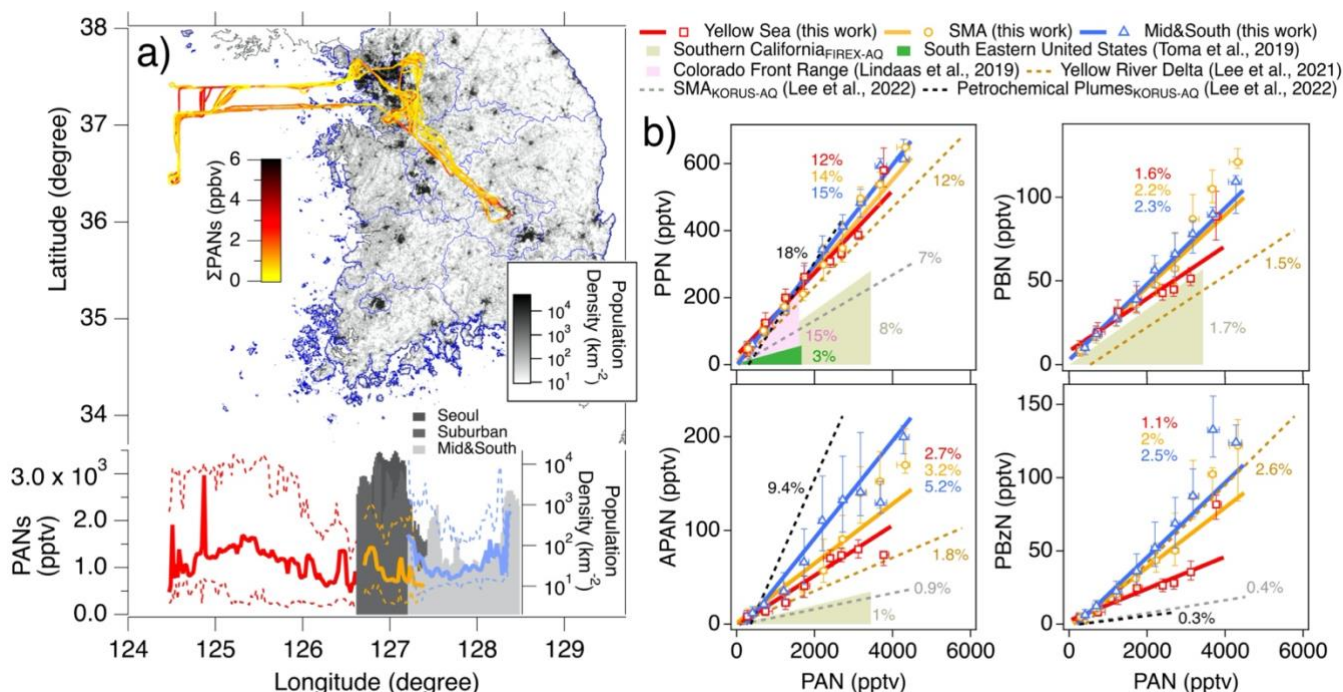


Figure 1. (a) (top) Map of South Korea DC-8 flight tracks colored by PANs mixing ratios, overlaid on a 1-km² population density map. (bottom) Longitudinal dependence of median PANs over the Yellow Sea (YS, red), Seoul Metropolitan Area (SMA, orange) and Mid and South (MS, blue) in 0.02° bins. The dashed lines represent longitudinal 25th and 75th percentiles of PANs. The shaded areas indicate median PD_{5-km} of the selected regions in 0.02° bins. (b) Scatter plots of median PAN homologues versus median PAN (i.e., homologue/PAN ratios), binned in intervals of 500-pptv PAN, with error bars denoting the 25th and 75th percentiles of the homologues in each bin. The red, orange, and blue solid lines indicate linear regression slopes for the 1s-merged data obtained over the YS, SMA and MS, respectively, during ASIA-AQ. Recent observations from Southern California (beige), the Southeastern United States (green), Colorado Front Range (pink), Yellow River Delta in China (light brown), and SMA (gray) and petrochemical plumes (black) from KORUS-AQ are also shown as dashed lines and shaded sectors.

High PANs levels were observed across the Korean Peninsula (mean $\pm 1\sigma$; 1.06 ± 0.64 ppbv) and the YS (1.43 ± 0.95 ppbv) during ASIA-AQ, with the maximum value (5.5 ppbv) measured in Gyeonggi Province within the SMA. These elevated PANs levels were accompanied by high levels of CH₂O, odd oxygen (O_x; approximated as O₃+NO₂), OA and aerosol concentrations (number, volume, and surface area) with generally $r^2 > 0.5$ across the selected regions (Fig. S5), indicating a strong photochemical relevance.

Correlations of PANs with aerosols were particularly strong during ASIA-AQ, exceeding those with O_x, similar to wintertime China (Qiu et al., 2019, 2020; Lu et al., 2019; Xu et al., 2021). During ASIA-AQ aerosol concentrations were measured by multiple instruments targeting different particle size ranges, providing additional constraints on these relationships. As shown in Fig. S5, aerosol concentrations (i.e., number, volume, and surface area) were highly correlated with PANs (r^2 up to ~ 0.9) with the strongest relationships for fine particles ($< 1 \mu\text{m}$) measured by an Ultra-High Sensitivity Aerosol Spectrometer (UHSAS). These observations suggest that PANs broadly capture the impacts of active VOC-NO_x



photochemistry on both gaseous and particulate pollution under cold conditions in East Asia, consistent with previous observations of the region (Lu et al., 2019; Xu et al., 2021; Lee et al., 2021).

250 Significant heterogeneity in both concentration and speciation of PANs demonstrated the influence of complex emission sources and photochemical processing. For example, the observed PANs distribution reflected the regional extent of photochemical pollution, extending well beyond source regions, and impacting downwind areas. Despite spanning more remote areas, median mixing ratios of PANs in the MS (990 pptv) and YS (1200 pptv) were higher than those in the SMA (840 pptv). This contrasts with the much higher population and industrial densities (e.g., ~40 times greater median PD_{5-km} and ID_{5-km}) in the SMA as well as more abundant precursor emissions. Figure 1a shows the spatial distribution, with the DC-8 flight tracks colored by PANs mixing ratios overlaid on a population density map (top panel), and longitudinal PANs quartiles and median PD_{5-km} in 0.02° bins (bottom panel). The longitudinal medians frequently exceeding 1 ppbv, including in remote areas, were particularly notable over the MS and YS. The longitudinal pattern of the MS showed increasing concentrations near the western and eastern boundaries, in close proximity to major industrial facilities including Banweol Industrial Complex (BIC), Daesan petrochemical complex (DPCC) and Gumi Industrial Complex (GIC), suggesting their potential contributions to pollutant distributions in the MS. In addition to South Korean sources, biomass burning plumes from China contributed to marked increases in PANs over the YS, as evidenced by concurrent enhancements of hydrogen cyanide (HCN) and acetonitrile (CH₃CN) (Figs. S5 and S6). These observations are consistent with high PAN episodes in Seoul (> 3 ppbv in 2019) (Savic et al., 2024) and at Gosan Climate Observatory (up to 2.4 ppbv), located near the southern YS (Han et al., 2017), attributed to long-range transport of biomass burning emissions from China. Consequently, the observed PANs distribution during ASIA-AQ is impacted by diverse sources including urban, industrial, and biomass burning emissions.

270 The speciation of PANs during ASIA-AQ exhibited heterogeneity as well. As shown in Fig. 1b, the relative abundance of PAN and its homologues (i.e., homologue/PAN ratios), derived from linear regression slopes, was at the upper end of previous reported values. These ratios exceeded those from recent U.S. observations from biogenic (Toma et al., 2019), industrial (Lindaas et al., 2019) and urban (Southern California during Fire Influence on Regional to Global Environments Experiment - Air Quality, FIREX-AQ 2019) regions, as well as during KORUS-AQ, and were most comparable to our measurements at a polluted remote site in China (Lee et al., 2021). Efficient thermal decomposition favors higher homologue/PAN ratios due to the ~15–25% faster thermolysis of PAN than its homologues (Kabir et al., 2014; Gomez et al., 2025). Consequently, the colder ASIA-AQ conditions in South Korea relative to the U.S. studies are unlikely to explain the higher ratios. Although not well constrained by experiments, potential slower OH oxidation of PAN compared to most larger homologues does not account for the observed homologue ratios and likely has a minor impact. However, it certainly contributes to a shorter lifetime of APAN (a few hours with $k_{\text{APAN}+\text{OH}} \sim 1 \text{ to } 2 \times 10^{-11} \text{ cm}^3 \text{ molecule}^{-1} \text{ s}^{-1}$ and $[\text{OH}] = 2 \times 10^6 \text{ molecules cm}^{-3}$) which depresses APAN/PAN ratios in cold environments. Instead, the elevated ratios point to a significant degree of photochemical pollution driven by photooxidation of complicated VOC mixtures, rarely observed outside East Asia in recent years.

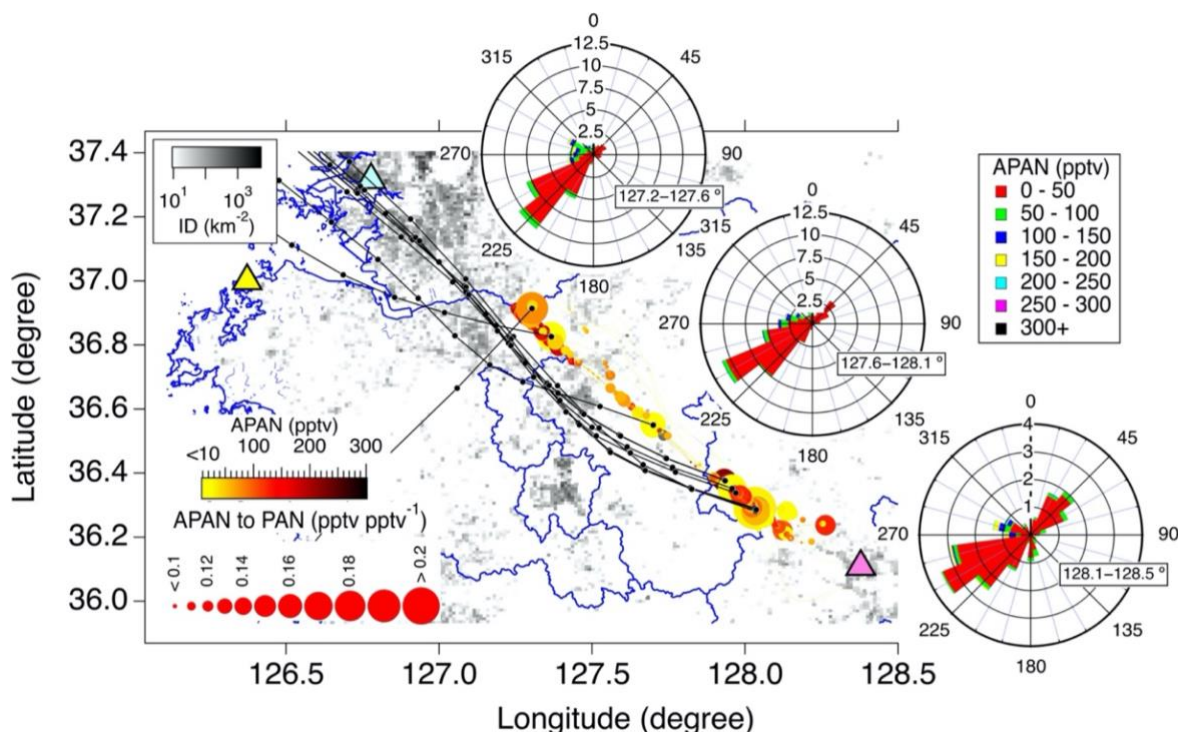


280 In addition, the variance in data points and the homologue ratios in Fig. 1b also point toward varying source contributions, with greater variability associated with homologues from more specific sources. Propionyl peroxyxynitrate (PPN) and sum of iso- and n-butyl peroxyxynitrate (PBN), formed from a broad range of anthropogenic alkanes and alkenes, showed relatively consistent ratios, whereas APAN and PBzN, formed from more reactive and source-specific alkenes and their product aldehydes (e.g., 1,3-butadiene and acrolein for APAN, and styrene and benzaldehyde for PBzN), showed considerable
285 variability. For example, PBzN/PAN ratios were lower over the YS compared to over the peninsula, consistent with significantly reduced aromatic concentrations (Fig. S6a). In contrast, this pattern is not observed for APAN/PAN ratios, despite the short lifetime of APAN due to removal by OH, because biomass burning is also a major source of APAN (Fig. S6b; Roberts et al., 2022).

Among all regions, the MS exhibited the highest homologue/PAN ratios, with the most pronounced enhancements
290 for the ratios of APAN and PBzN. A consistent increase in the ratios for APAN and PBzN was observed in the MS with their concentrations reaching hundreds of pptv and the strongest regional correlation ($r^2 \sim 0.5$) between the two species. Based on our previous field measurements in various environments, such elevated levels and ratios have only been observed in regions with dominant contributions of petrochemical and biomass burning emissions. While PANs observations are widely used to assess VOC–NO_x photochemistry over the past decades, their application to APAN and PBzN remain scarce, particularly in
295 East Asia. Given their clear enhancements, limited precursor sources and low background concentrations, APAN and PBzN can provide valuable insights into complex photochemical processes. Therefore, in the following section, we present a case study using these tracers to investigate regional-scale pollution in the MS.

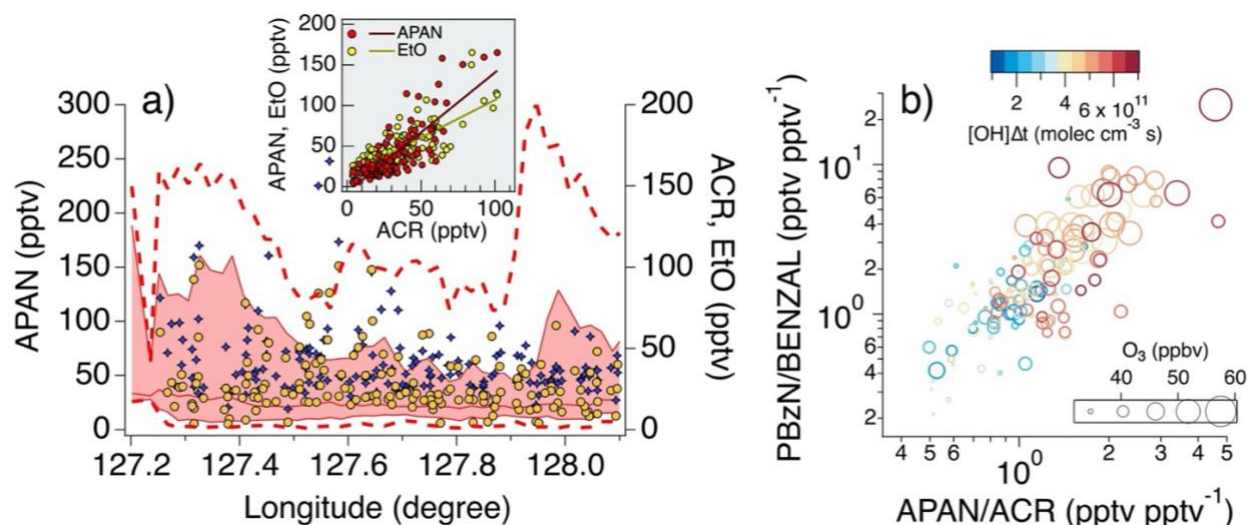


3.1 Investigation of MS pollution using APAN and PBzN



300 **Figure 2.** APAN and APAN/PAN ratios over the MS illustrated as marker color and size, respectively, overlaid on a 1-km² industrial facility density map, with wind rose plots for three 0.5° longitudinal bins colored by APAN mixing ratios. The black lines with solid circles indicate back trajectories initiated at the 10 largest APAN/PAN values. The cyan, yellow, and magenta triangles indicate the locations of Banweol Industrial, Daesan Petrochemical, and Gumi Industrial Complexes, respectively.

We investigate recent photochemical processes and regional pollution over the MS. Although significant pollution
305 was observed during ASIA-AQ, this region is understudied. For this purpose, we utilize APAN as a primary tracer, complemented by PBzN as an additional proxy for anthropogenic aromatic sources. APAN is particularly useful due to its short atmospheric lifetime and substantial enhancements over low background levels observed in this region. Figure 2 presents spatial distributions of APAN and APAN/PAN ratios, along with wind rose plots in 0.5° longitudinal bins and back trajectories (HYSPLIT; Stein et al., 2015) initiated at the 10 largest APAN/PAN values. Elevated APAN concentrations and APAN/PAN
310 ratios up to 343 pptv and 0.2, respectively, were consistently associated with west and northwesterly winds (270°–315°), indicating transport from west coast industrial sources. Although median APAN levels were low (24 pptv), upper-end values were elevated and spatially extensive. Longitudinal profiles in Fig. 3a show APAN increasing near the eastern and western boundaries, consistent with the PANs distribution in Fig. 1b. Additional enhancements in the southeast were associated with northeasterly winds, suggesting local influence from the Gumi Industrial Complex.



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Figure 3. (a) APAN, acrolein (ACR) and EtO (ethylene oxide) as a function of longitude over the MS. The solid red line and shaded area indicate longitudinal 50th, 25th, and 75th percentiles, respectively, of APAN in 0.02° bins, and the dashed red lines are the longitudinal 1st and 99th percentiles. The orange and blue markers represent ACR and EtO mixing ratios. The inset shows correlation of ACR with APAN (red markers with black linear fit line) and with EtO (yellow markers with beige linear fit line). (b) Scatter plots of PBzN/benzaldehyde (BENZAL) versus APAN/ACR over the MS. The markers are colored and sized by OH exposure and O₃ mixing ratios, respectively.

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The inset of Fig. 3a shows acrolein (ACR), a primary aldehyde precursor of APAN, correlated most strongly ($r^2 > 0.6$) with APAN and ethylene oxide (EtO) among all ASIA-AQ measurements. It is noted that concurrent airborne measurements of these species were conducted for the first time in East Asia. Given the short lifetimes of APAN and acrolein, the enhancements are indicative of recent emissions from local to regional sources. The source of APAN and acrolein being dominated by biomass burning in the MS is unlikely due to a lack of correlation with HCN and CH₃CN. The positive relationship between acrolein and benzene ($r^2 \sim 0.6$) measured by TOGA in the SMA, but not in the MS, suggests that urban emissions do not explain the observations over the MS. Instead, the results point to industrial sources, particularly petrochemical facilities along the west coast, as the origin of both acrolein and APAN, supported by strong correlations with EtO, a tracer almost exclusively emitted from petrochemical activity (Robinson et al., 2024).

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Recently, Robinson et al. (2024) reported EtO levels reaching hundreds of ppTV in southeastern Louisiana's petrochemical corridor, exceeding EPA estimates (median difference of 21 ppTV) and highlighting both the scarcity of ambient EtO data and potential negative health risks. Further evidence for a similar source attribution in South Korea is provided by back trajectories in Fig. 2, which intersects the BIC, a major industrial complex hosting roughly 13,000 metal manufacturing and 1400 petrochemical facilities, including synthetic rubber production (Chae et al., 2024). Although the back trajectories of the selected points did not directly intersect the DPCC, observations during KORUS-AQ demonstrated that its emissions influence the YS and southern Chungcheong regions adjacent to the MS. For example, our previous work showed efficient production of APAN and acrolein from 1,3-butadiene oxidation in plumes near DPCC and over the YS (Lee et al., 2022). In addition, the highest acrolein level observed during ASIA-AQ over the YS (600 ppTV) coincided with the maximum 1,3-



340 butadiene mixing ratio (1447 pptv) and elevated APAN (90 pptv) during ASIA-AQ, indicating the continued impact of petrochemical emissions in South Korea and their potential contribution to observations over the MS. Given the rapid formation of acrolein from 1,3-butadiene oxidation, direct source attribution is challenging. However, the strong positive relationship of APAN with EtO supports a dominant contribution from petrochemical emissions and suggest elevated exposure to highly toxic pollutants in the MS.

345 Previous model simulations by Lee et al. (2022) predict PBzN formation from styrene oxidation in the DPCC plumes. Our PBzN measurements during ASIA-AQ showed consistent enhancements alongside APAN, in agreement with the simulations. Figure 2c shows both APAN/acrolein and PBzN/benzaldehyde ratios increase generally with OH exposure, which provides a proxy for photochemical processing, consistent with secondary formation from aldehyde precursors. The OH exposure, derived using Eq. (S1) based on observed isopropyl nitrate to propane ratios (see Sect. S5.1) captured the expected
350 decay in the NO_x contribution to NO_y (approximated as $\text{NO}_x + \text{PANs} + \text{HNO}_3 + \text{pNO}_3^-$) and the increase in PANs to aldehyde ratios with increasing OH exposure (Fig. S7). Both APAN/acrolein and PBzN/benzaldehyde ratios increased with O_3 as well, further implicating photochemical oxidation of petrochemical VOCs in O_3 production, consistent with markedly significant O_3 levels up to 250 ppbv in petrochemical plumes during KORUS-AQ (Fried et al., 2020; Cho et al., 2021; Lee et al., 2022). Together, observations over the MS provide strong observational evidence of regional-scale photochemical pollution
355 influenced by petrochemical emissions, with APAN, PBzN and their precursors as effective tracers of this impact.

4 Observation-based kinetic calculations and box modeling for diagnosis of VOC– NO_x photochemistry

4.1 Instantaneous production of PANs

To interpret the observed distributions of PANs, we use a combined analysis based on kinetic calculations and model simulations to estimate instantaneous production rates of PANs and their precursor contributions. For this purpose, we
360 investigate the influence of precursor oxidation on PANs formation by focusing on the relative abundances of speciated PANs, which are less sensitive to atmospheric mixing and dilution than absolute concentrations. These processes are difficult to constrain in model simulations, particularly in regions with complex and uncertain emissions. Figure 4a compares observed fractional contributions with modeled instantaneous production rates ($P(\text{PANs})$), where $P(\text{PANs})$ denotes the combined production of PANs and corresponding peroxyacyl (PAs) radicals as a chemical family. The agreement between observations
365 and $P(\text{PANs})$ suggests that local photooxidation is consistent with the chemical evolution responsible for the observed PANs distribution. This agreement was strongest over inland regions, where the influence of local emissions and recent photochemistry is expected to be more pronounced.

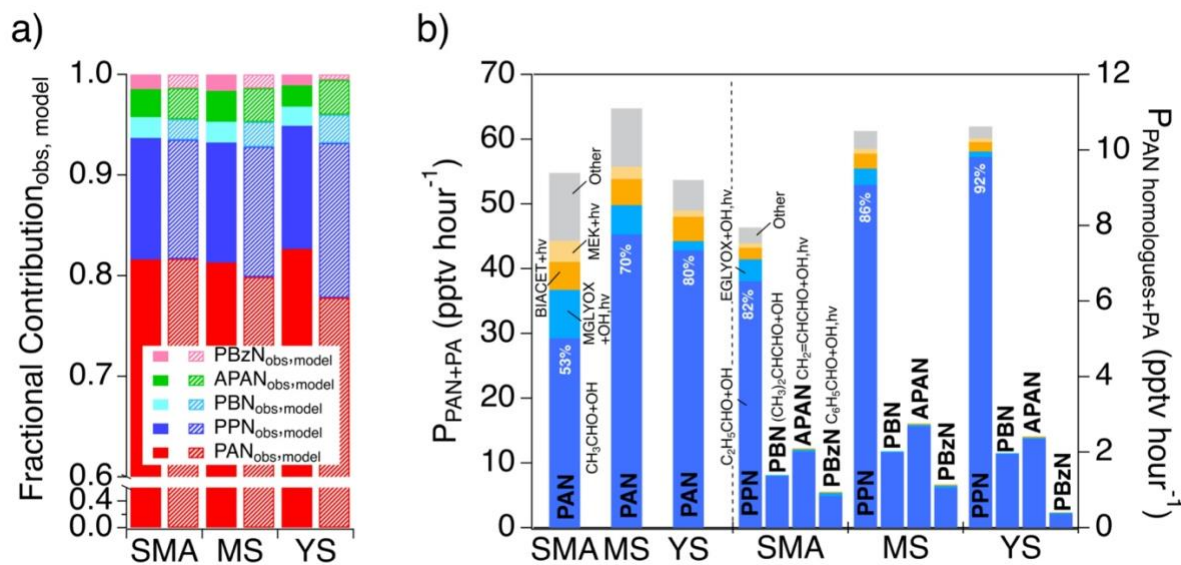


Figure 4. (a) Averaged fractional contributions of observed PANs compounds (bright colored bars) and those of modeled instantaneous production rates of PANs, $P(\text{PANs})$, (colored bars with patterns) over the SMA, MS, and YS. (b) Averaged $P(\text{PANs})$ over the SMA, MS, and YS along with precursor contributions. The left three bars correspond to $P(\text{PAN})$ and other bars corresponds to $P(\text{PAN homologues})$.

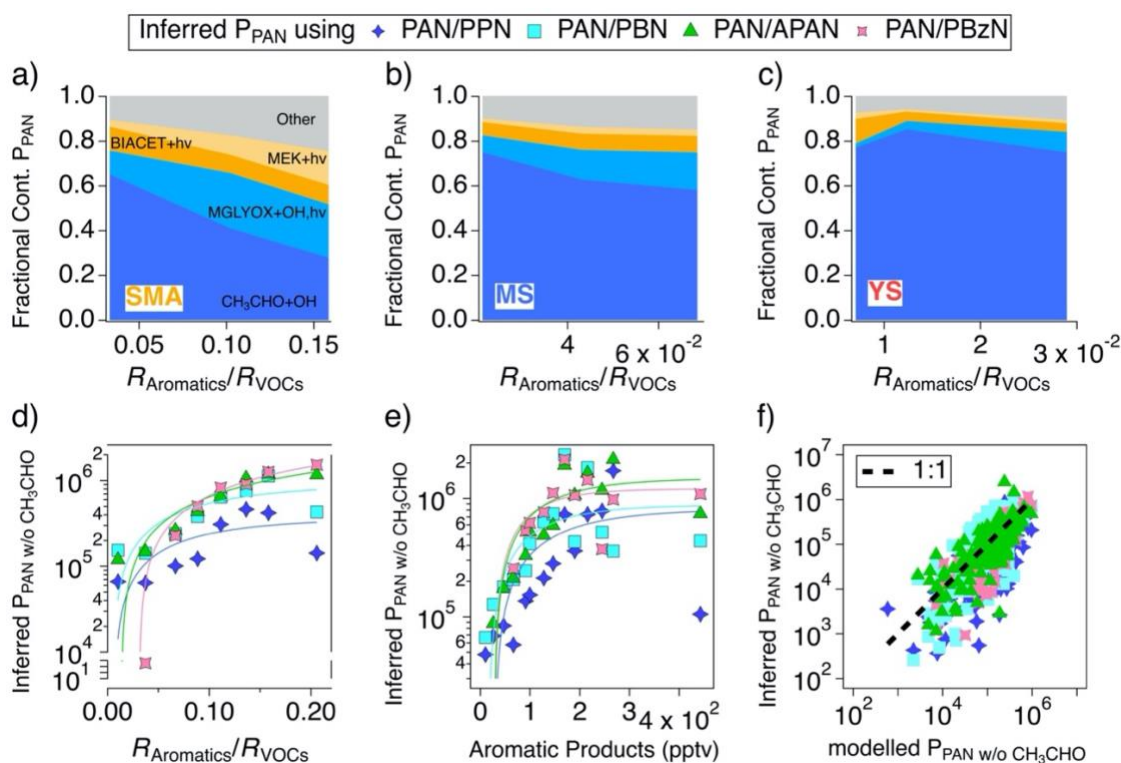
Regional differences in $P(\text{PANs})$ and precursor contributions are illustrated in Fig. 4b. $P(\text{PANs})$ was higher over the MS and YS, consistent with elevated PANs mixing ratios and indicative of abundant precursors and active local formation. In contrast to regions with simpler VOC composition, in which acetaldehyde often dominates PAN production, our analysis illustrates non-acetaldehyde sources including methyl ethyl ketone (MEK), biacetyl (BIACET) and methyl glyoxal (MGLYOX), accounted for up to 47% of PAN formation on average. While MEK and BIACET were measured, MGLYOX from model simulations was used for $P(\text{PANs})$ calculations. The findings in this work suggest that less common precursors are critical for PAN chemistry over South Korea, which has been hypothesized in previous studies from KORUS-AQ (Nault et al., 2024) and from other polluted environments (Liu et al., 2010).

As the structure of PANs becomes more complex, the range of possible precursors narrows (Roberts et al., 2007). Consistent with this general tendency, contribution of unconventional precursors on PAN homologue formation was limited. Although PPN had a non-negligible contribution from ethyl glyoxal, analogous with MGLYOX contribution for PAN formation, propanal dominated its production. Larger homologues including APAN, PBN and PBzN were produced almost exclusively from their respective aldehyde precursors. Though present at much lower levels than PAN, the simpler formation pathways of the larger PAN compounds make them useful photochemical tracers for their precursor sources.

Constraining PAN formation from non-acetaldehyde precursors, hereafter $P(\text{PAN})_{\text{w/o } \text{CH}_3\text{CHO}}$, has been a focus of research, particularly in environments impacted by isoprene and aromatics (Roberts et al., 2006; Liu et al., 2010; Nault et al., 2024; Travis et al., 2024). Major PAN precursors identified in the previous section such as MEK, BIACET, and MGLYOX can originate from both biogenic and anthropogenic sources (Fischer et al., 2014), complicating source attribution in East Asia, including South Korea, where both sources are prominent (Simpson et al., 2019). With limited isoprene during wintertime

ASIA-AQ conditions, PAN formation is expected to be dominated by anthropogenic sources, providing a testbed to access their impacts on PAN precursors.

Given that MGLYOX is expected to contribute significantly to $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ and is a known product of aromatic oxidation (e.g., Fu et al., 2008; Nishino et al., 2010), we investigate the fractional contribution of PAN precursors as a function of aromatic contribution to OH reactivity (Fig. 5a–c). The modeled $P(\text{PAN})$ exhibited decreasing relative contribution from acetaldehyde with increasing aromatic contribution to OH reactivity. This is most prominent over the SMA, where aromatic contributions are the highest (Fig. 5a). Over the SMA, the acetaldehyde contribution decreased to ~30%, with substantial contributions from MEK, BIACET, and MGLYOX, consistent with regional mean $P(\text{PAN})$ shown in Fig. 4. The contribution of MEK was also increased in regions with greater aromatic influence, while the impacts of BIACET are relatively constant.



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Figure 5. (a–c) Modeled fractional contributions from PAN production channels as a function of aromatic contribution to OH reactivity from VOCs, $R_{\text{Aromatics}}/R_{\text{VOCs}}$, over the SMA, MS, and YS. Inferred $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ as a function of (d) $R_{\text{Aromatics}}/R_{\text{VOCs}}$ and (e) mixing ratios of aromatic oxidation products. The solid fit lines in (d) and (e) are provided as a visual aid. (f) Inferred $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ based on kinetic calculations compared to modeled $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$. The bold dashed line is the 1:1 line. Inferred and modeled $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ are in units of molecules $\text{cm}^{-3} \text{s}^{-1}$.

To complement our model analysis, we use kinetic calculations to infer $P(\text{PAN})$ from non-acetaldehyde sources (inferred $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$), using Eqs. (S2)–(S3) in Sect. S2.2. This analysis provides an additional constraint on $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ to our model analysis because direct experimental quantification remains challenging due in part to significant interferences in even state-of-the-art mass spectrometric and optical techniques for MGLYOX measurements (Koss et al.,



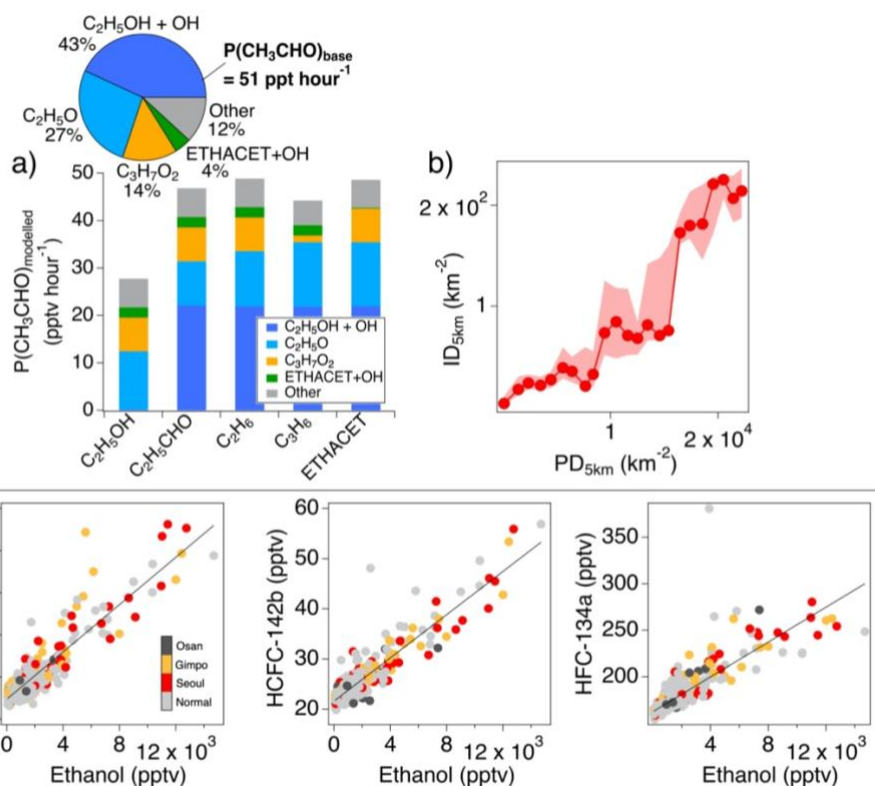
410 2018). Briefly, the method leverages extensive measurements of PANs and their precursors and is based on two assumptions:
(1) observed PAN-to-homologue ratios are representative of the ratios of their instantaneous production rates; and (2) PAN
homologues are dominantly formed from their corresponding aldehydes. These assumptions were assessed in the previous
section using model simulations and found reasonable.

The inferred $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ was illustrated as a function of aromatic contribution to OH reactivity (Fig. 5d) as well
415 as the summed concentrations of observed aromatic oxidation products including benzaldehyde, cresol, phenol, nitroresol,
and nitrophenol (Fig. 5e). The inferred $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$ increased with these proxies for aromatic oxidation, consistent with
the modeled contribution of MGLYOX in this work and previous model studies in East Asia (Liu et al., 2010; Fischer et al.,
2014). Notably, the values based on PAN/PPN ratios were generally lower than those from other homologues, consistent with
420 minor contributions to PPN from non-propanal precursors (8–18%) and almost exclusive contribution of the respective
aldehydes to larger PAN homologues. When compared to the model-derived $P(\text{PAN})_{\text{w/o CH}_3\text{CHO}}$, the kinetic estimates were in
reasonable accord, supporting the validity of the estimations from two different methods (Fig. 5f). Overall, our observation-
based diagnostic, combining model simulations and kinetic calculations, provides strong evidence for the contribution of non-
acetaldehyde sources to PAN formation in South Korea. This contribution is particularly important in the SMA where
aromatics are more abundant.

425 4.2 Contribution of ethanol and other precursors to secondary acetaldehyde production

Despite the important contributions of non-acetaldehyde sources to PAN production, acetaldehyde remains the most
important precursor of PAN on average. In urban environments, acetaldehyde is often dominated by secondary production
from oxidation of hydrocarbons from anthropogenic emissions, although acetaldehyde can originate from primary emissions
from various anthropogenic and biogenic sources (e.g., Grosjean et al., 2002; Millet et al., 2010; de Gouw et al., 2018). Here,
430 we investigate the source of acetaldehyde as it is of particular importance for PAN as well as O₃ formation (e.g., Sect. 3.2.3)
in South Korea.

The pie chart in Fig. 6a shows fractional contribution of direct precursors of modeled acetaldehyde production,
 $P(\text{CH}_3\text{CHO})$, with a mean value of 51 ± 46 (1σ) pptv h⁻¹ over South Korea. The bar charts illustrate how $P(\text{CH}_3\text{CHO})$ and its
direct precursors change upon removal of the selected major precursors in model simulations (i.e., zero-out sensitivity analysis)
435 for South Korea. The most significant reduction of $P(\text{CH}_3\text{CHO})$ was found when ethanol was set to zero. Compared to the
base case simulation, $P(\text{CH}_3\text{CHO})$ decreased by nearly a half. This result from a simple diagnostic box modeling is consistent
with a 50% reduction in acetaldehyde without including ethanol in GEOS-Chem transport model simulations for the SMA
during KORUS-AQ (Travis et al., 2024). To a lesser extent, oxidation of more common hydrocarbon precursors (e.g., ethane
and propene) and carbonyls (e.g., propanal and ethyl acetate) also contributed to $P(\text{CH}_3\text{CHO})$.



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Figure 6. (a) Pie chart illustrating the fractional contributions to direct acetaldehyde production, $P(\text{CH}_3\text{CHO})$, from individual pathways in the base-case model simulation (mean $P(\text{CH}_3\text{CHO}) = 51 \text{ ppt h}^{-1}$). $\text{C}_2\text{H}_5\text{OH} + \text{OH}$ represent ethanol oxidation by OH. $\text{C}_2\text{H}_5\text{O}$ and $\text{C}_3\text{H}_7\text{O}_2$ denote alkoxy and hydroxy alkoxy radicals, respectively, which decompose to form acetaldehyde. ETHACET + OH indicates OH oxidation of ethyl acetate. Others include minor pathways such as OH oxidation of MEK and PPN, and ozonolysis of propene and butene isomers. The bar charts illustrate changes of $P(\text{CH}_3\text{CHO})$ in response to removal of the selected major precursors of acetaldehyde in model simulations, so-called "zero-out sensitivity tests." (b) Scatter plot showing median industrial facility density ($\text{ID}_{5\text{-km}}$) as a function of population density ($\text{PD}_{5\text{-km}}$), binned in intervals of 1000, with shaded areas indicating the 25th and 75th percentiles in each bin. (c) Scatter plots of CHBrCl_2 , HCFC-142b and HFC-134a versus ethanol, with colored and gray markers indicating measurements during low approaches and normal operations, respectively.

The substantial contribution of ethanol to $P(\text{CH}_3\text{CHO})$, particularly due to its high abundance at low altitudes (median = 2 ppbv, maximum = 13 ppbv at altitudes < 500 m), underscores its importance for PAN and motivate further investigation on its local sources in South Korea. Recent model analyses based on KORUS-AQ data have identified ethanol as a key precursor to acetaldehyde and PAN, consistent with findings in this work, using estimated ethanol levels from ethanol/methanol ratios of 0.4 obtained from previous ground-based campaign (the Megacity Air Pollution Studies-Seoul; MAPS-Seoul) (Nault et al., 2024; Travis et al., 2024). The ASIA-AQ observations over South Korea showed a moderately correlated relationship between ethanol and methanol ($r^2 = 0.44$) and an ethanol/methanol ratio of 0.48, derived from the slope of a linear regression, supporting these estimates. In addition, Travis et al. (2024) demonstrated that increasing ethanol emission factors by a factor of 40 over those in current emission inventory (KORUS-AQ v5) was necessary to reduce model bias in PAN over the SMA from -50% to -23%, while the significant increase in ethanol was attributed to VCPs.



460 Consistent with this source attribution, Beaudry et al. (2025) reported that reproducing ethanol levels during KORUS-
AQ and MAPS-Seoul campaigns required per capita VCP emissions in South Korea to be 2.4 times higher than in the U.S. In
contrast to the implementation of population-based emissions of ethanol from VCPs, no significant relationship was found
between ethanol and PD_{5km} , suggesting ethanol is from a source unrelated to local population density or is distributed over
large areas. In addition, Fig. 6b illustrates co-location of industrial facilities and populated areas using scatter plot of ID_{5km}
465 over PD_{5km} . This spatial overlap of the sources may partially explain improved model performance implementing population-
based emissions when ethanol emissions from industrial sources contribute to the observed distribution. This indicated that a
comprehensive source attribution including VCP emissions from industrial and consumer products, as well as other possible
sources (e.g., fuel combustion), is critical for accurately assessing atmospheric impacts of ethanol. Supporting this
interpretation, Fig. 6c shows that ethanol correlated most strongly with halocarbons, including bromodichloromethane
470 ($CHBrCl_2$), HCFC-142b, and HFC-134a with $r^2 > 0.8$. The strong positive relationships were particularly noticeable during
low approaches near major urban airports (i.e., Seoul and Gimpo), suggesting surface emissions from industrial (e.g., chemical
production, refrigerants and blowing agents) and solvent-related sources (ATSDR, 2005, 2020 and references therein). Beyond
air quality implications, investigation of ethanol and halocarbon emissions over South Korea has the potential to improve
understanding of the sources and distributions of short-lived (O_3) and longer-lived (HCFCs, HFCs) climate forcers, given
475 increasing background levels of such halocarbons with unidentified sources in East Asia (Choi et al., 2024).

In summary, ethanol and other species were significant sources of acetaldehyde. Previous studies have attributed
ethanol emissions to VCPs based on findings from U.S. cities. However, our results indicate that such attribution requires a
more detailed understanding of local sources, including industrial activities near populated areas. The heterogenous distribution
of VOCs and PANs observed in this work support this conclusion.

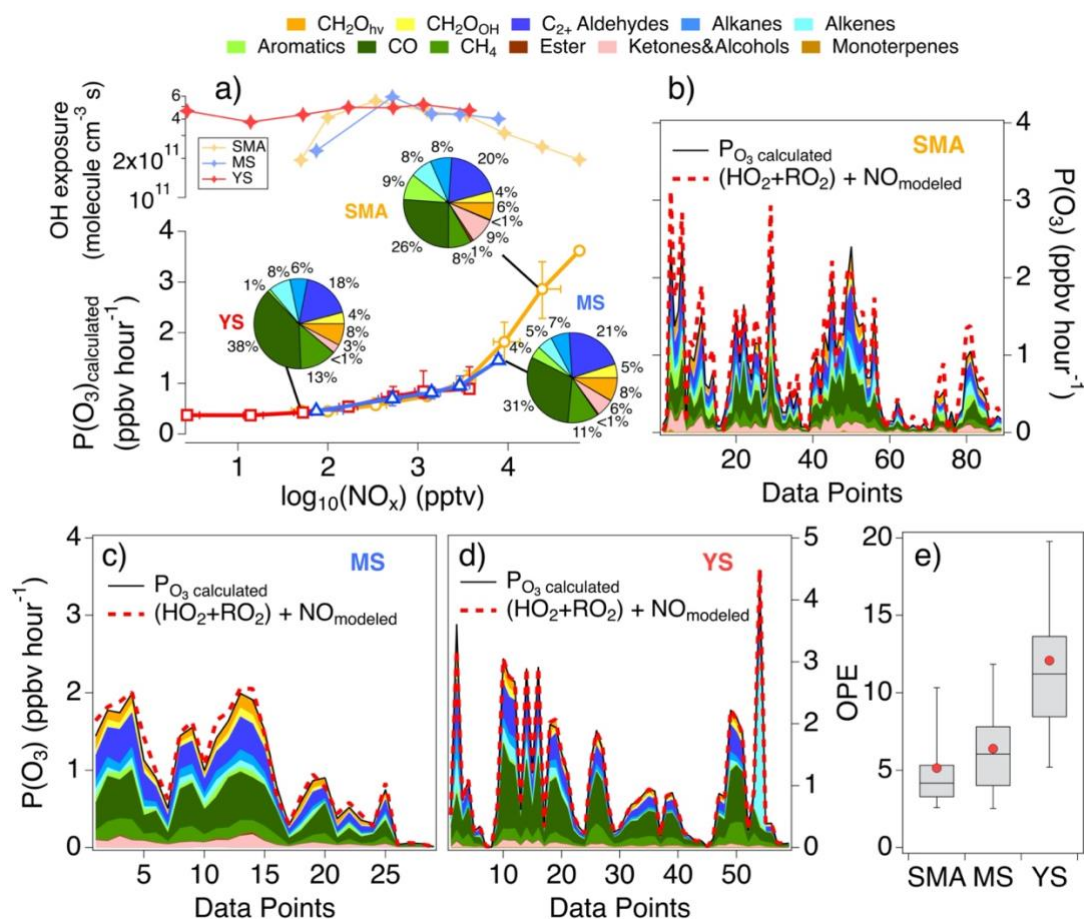
480 4.3 Instantaneous production rates, efficiency, and chemical regimes of O_3

Ozone formation is evaluated based on instantaneous ozone production rates, $P(O_3)$, and efficiency, OPE. For this
evaluation we use a combined analysis incorporating kinetic calculations and model simulations. This analysis enables a
detailed diagnostic of ozone production by integrating observation-based estimates using experimental parameters with near-
explicit chemical modeling that includes precursors not directly measured such as radicals. Figure 7a shows $P(O_3)$ from kinetic
485 calculations using Eq. (3) and OH concentration of 2×10^6 molecules cm^{-3} as a function of NO_x on a logarithmic scale, with
OH exposure shown in the upper panel. The general increase in $P(O_3)$ is due to increasing OH reactivity with NO_x , as the
calculation does not account for suppression of OH at high NO_x levels.

$P(O_3)$ from kinetic calculations is more effective for evaluating relative precursor contributions independent of OH.
The pie charts in Fig. 7a show substantial contributions of C_{2+} aldehydes to $P(O_3)$, second in importance to CO. Other important
490 contributors include formaldehyde as well as ketones and alcohols such as MEK, methanol, ethanol, and isopropanol, which
together account for ~95% of this category. Less significantly, primary hydrocarbons including methane (CH_4), alkanes,
alkenes, and aromatics over South Korea showed similar contributions, while aromatic impacts were minimal in the YS. Less-



reactive species such as CO and CH₄ were more important over the YS, likely due to sampling of air masses with biomass burning influence and with longer time since emissions.



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Figure 7. (a) Median ozone production, $P(O_3)$, from kinetic calculations using $[OH] = 2 \times 10^6$ molecules cm^{-3} for the SMA (orange line with circles), MS (blue line with triangles) and YS (red line with squares) in 0.5-logarithmic NO_x bins. Vertical and horizontal lines represent 25th and 75th percentiles on the respective axes. The OH exposure (lines with markers) for each region is shown in the upper panel. Pie charts indicate the fractional contribution of each of VOC classes (defined in the legend) to $P(O_3)$. (b–d) Comparisons of $P(O_3)$ from kinetic calculations using modeled $[OH]$ (black lines with shaded areas representing precursor contributions) and model simulations based on the sum of reactions of peroxy radicals with NO over the selected regions (red dashed lines). Data points on the x-axis represent sequential indices of model output, used to illustrate the comparison between kinetic calculations and model simulations. (e) Box and whisker plot of instantaneous ozone production efficiency (OPE), along with regional averages (red markers).

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When modeled OH was used in the kinetic calculations, instead of using a constant value for generalization, $P(O_3)$ from kinetic calculations was in excellent agreement with model simulations based on Eq. (2) (Fig. 7 b–d). This indicates that the modeled oxidative chemistry driving ozone production and radical cycling can be explained by the current understanding of observed VOC photooxidation. Consistent with the averaged impacts shown in Fig. 7a, oxygenates including C₂₊ aldehydes and formaldehyde showed significant and persistent contribution in all regions. In addition, episodic enhancements of alkenes dominating $P(O_3)$ was observed in the YS and attributed to sampling of relatively fresh petrochemical plumes, also described



510 in Sect. 3.1. Another important measure for ozone production is OPE. Figure 7e shows modeled OPE using box plots, with
 515 red markers indicating regional averages. OPE generally decreased with increasing NO_x , resulting in the lowest and highest
 values in the SMA and YS, respectively. Higher OPE indicates more efficient radical propagation relative to termination by
 NO_x . The interquartile ranges of OPE for the SMA (3.3–5.3) and MS (4–7.8) were comparable to the previous reported values
 in East Asia ($< \sim 10$) in winter–spring, though only limited values are available for winter (Lin et al., 2011; Lee et al., 2021;
 Kim et al., 2020; Oak et al., 2019).

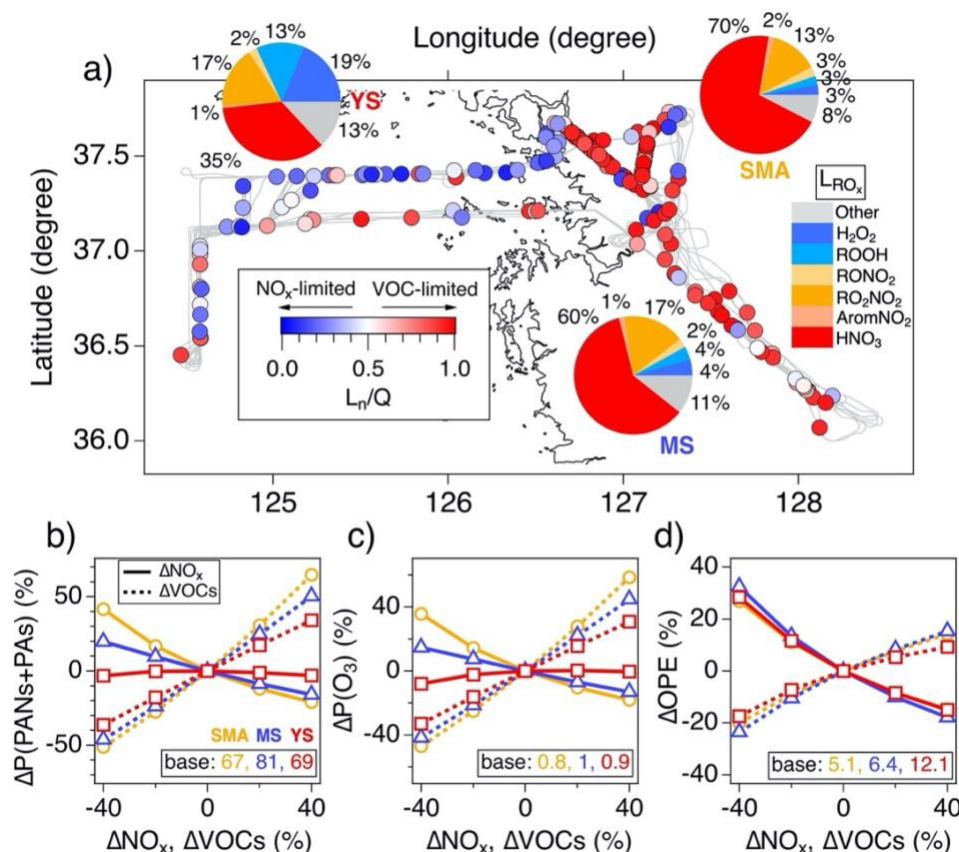


Figure 8. (a) Averaged fractional contributions of modeled instantaneous radical loss pathways over the SMA, MS, and YS (pie charts), along with markers colored by instantaneous fraction of radical loss by NO_x in total loss (L_n/Q). Model sensitivity of (b) $P(\text{PANs})$, (c) $P(\text{O}_3)$ and (d) OPE to perturbations of NO_x and VOCs.

520 A key factor determining OPE and the prevailing chemical regime is the balance of radical ($\text{RO}_x = \text{OH} + \text{HO}_2 + \text{RO}_2$)
 termination pathways, including self-reactions of peroxy radicals and reactions with NO_x , which removes radicals and NO_x
 from ozone producing photochemical cycles. The modeled instantaneous radical loss indicates that radical loss by reaction
 with NO_x , producing nitric acid (HNO_3) and PANs, dominates over the SMA (88%) and MS (80%) (Fig. 8a). Such dominant
 contributions of HNO_3 and PANs to radical loss are consistent with wintertime observations in Beijing based on direct radical
 525 measurements (Tan et al., 2018; Lu et al., 2019). Minor radical termination pathways include formation of peroxides, alkyl



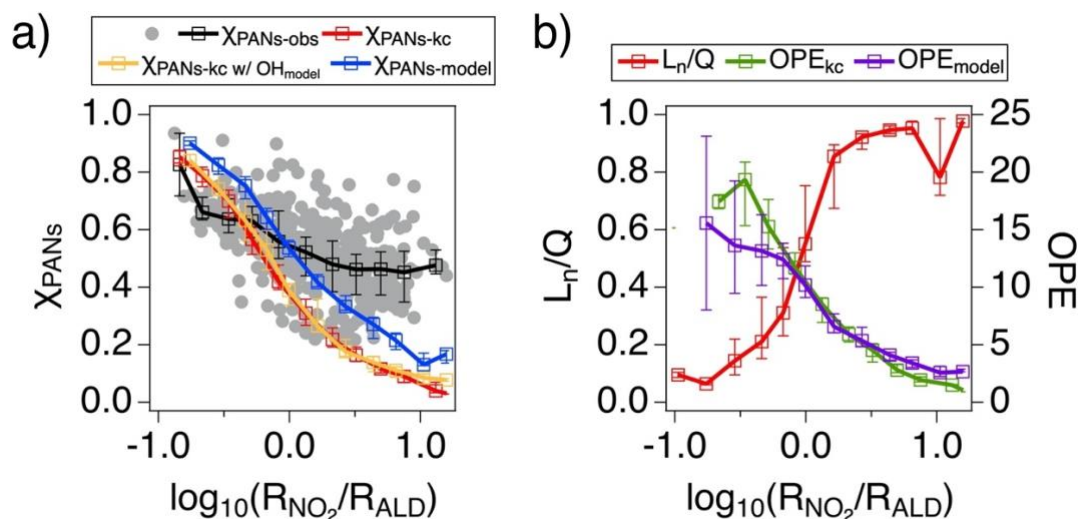
nitrates, HONO, pernitric acid (HO_2NO_2) and nitroaromatic compounds, though the chemistry of nitroaromatics in the MCM mechanism may be uncertain and needs further investigation (Bates et al., 2021; MacFarlane et al., 2025).

In addition to OPE, the ratio of radical loss by NO_x (L_n) to total loss (Q), L_n/Q ratio, provide an additional diagnostic measure for ozone formation regimes, with $L_n/Q > 0.5$ indicating VOC-limited regimes and $L_n/Q < 0.5$ indicating NO_x -limited regimes (Kleinman et al., 2002). Figure 8a shows that median L_n/Q values of 0.9 over South Korea, representative of VOC-limited regimes with major radical loss driven by HNO_3 and PANs formation in the region. Over the YS, radical-radical reactions producing peroxides become increasingly important, resulting in a lower median L_n/Q of 0.4.

To investigate potential impacts of emission changes on chemical regimes, model sensitivity simulations were conducted by evaluating $P(\text{PANs})$, $P(\text{O}_3)$ and OPE responses to perturbations of NO_x and VOCs constraints ($\pm 40\%$ of the base case). Figure 8b–c show the average relative changes of $P(\text{PANs})$ and $P(\text{O}_3)$ exhibit a strong positive response to VOCs across all regions, while NO_x perturbation produces opposite responses in South Korea and a weaker effect in the YS. In contrast to this regional difference, OPE showed more uniform response across regions as it represents the average changes in the ratio between formation of radicals and NO_x oxidation products. The model sensitivity results point out that PANs and ozone formation during ASIA-AQ occurred in VOC-limited conditions and suggest that VOC reductions would be more effective than NO_x controls for mitigation of photochemical pollution.

4.4 Diagnostic kinetic and model analysis on NO_x loss pathways

A defining feature of the ASIA-AQ observations was the marked heterogeneity in pollutant distributions, reflecting a broad range of photochemical conditions and pollution intensities (e.g., NO_x levels spanning orders of magnitude; Fig. 7a). While NO_x removal was dominated by HNO_3 and PANs, which together accounted for $\sim 90\%$ of modeled NO_x loss over South Korea, their relative contributions varied largely during the campaign, in part due to this heterogeneity. Romer et al. (2020) evaluated observed relative contributions of HNO_3 and alkyl nitrates (RONO_2) to NO_x loss during the summer daytime across multiple campaigns in the U.S. over 15 years, using kinetic calculations based on instantaneous production rates. They pointed out the relative contribution of HNO_3 decline with the OH reactivity ratio of NO_2 to the sum of VOCs ($R_{\text{NO}_2}/R_{\text{VOCs}}$), suggesting increased importance of RONO_2 chemistry with decreases in NO_x levels.



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Figure 9. (a) Median observed fractional contribution of PANs to the total NO_x loss based on observations ($\chi_{PANs-obs}$; black line with markers) along with median ratios of instantaneous PANs production rates to the combined rates of PANs and HNO_3 production from kinetic calculations using $[OH] = 2 \times 10^6$ molecules cm^{-3} ($\chi_{PANs-kc}$; red line with markers) and modeled $[OH]$ ($\chi_{PANs-kc}$ w/ OH_{model} ; orange line with markers), and from model simulations ($\chi_{PANs-model}$; blue line with markers) in 0.2-logarithmic R_{NO_2}/R_{ALD} bins. All data points used to calculate $\chi_{PANs-obs}$ are illustrated as gray markers. (b) Median modeled L_n/Q (red line with markers) and OPE (OPE_{model} ; purple line with markers) and kinetic calculated OPE (OPE_{kc} ; green line with markers) in 0.2-logarithmic R_{NO_2}/R_{ALD} bins. The error bars in both figures indicate 25th and 75th percentiles in each bin.

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Using a similar kinetic approach to Romer et al. (2018), we investigate the observed fractional contribution of PANs to total NO_x loss ($\chi_{PANs-obs}$), defined here as the sum of observed PANs, HNO_3 , and particulate inorganic nitrate (pNO_3^- inorganic). pNO_3^- inorganic was estimated by excluding organic nitrate contribution (~10% of pNO_3^- over South Korea) from total nitrate measured by the University of Colorado Aerosol Mass Spectrometer (CU aircraft AMS), following the method of Day et al. (2022). This diagnostic analysis may be particularly relevant for wintertime East Asia, where cold conditions extend PANs lifetime (effective lifetime of ~ 3 days) and enable efficient NO_x sequestration (e.g., Fig. 8). The competition among HNO_3 and PANs formation is a critical factor that controls NO_x lifetimes and the spatial extent of ozone pollution, as HNO_3 formation constitutes a permanent sink, whereas PANs serve as a reservoir and can facilitate NO_x transport on local–regional scales.

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During ASIA-AQ, $\chi_{PANs-obs}$ varied substantially (0.2–0.94) and had an inverse relationship with the OH reactivity ratio of NO_2 to the sum of observed aldehyde precursors of PANs (R_{NO_2}/R_{ALD}). The decrease in $\chi_{PANs-obs}$ with R_{NO_2}/R_{ALD} suggests the importance of competition between OH reacting with NO_2 to form HNO_3 versus with aldehydes to form PAs, which promotes PANs formation, in determining NO_x loss pathways. It should be noted that R_{ALD} is more relevant than R_{VOCs} used in previous studies for the analysis herein and may provide a more consistent basis for future analyses, given that kinetic calculations are sensitive to the selection of VOCs and their associated kinetic parameters (e.g., alkyl nitrate branching ratios and yields).

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The observed dependence of $\chi_{PANs-obs}$ with R_{NO_2}/R_{ALD} is consistent with kinetic expressions (Eq. (S4–S6)) that
575 define the ratio of instantaneous PANs production rates to the combined rates of PANs and HNO_3 production. These equations
exhibit a negative relationship of χ_{PANs} with R_{NO_2}/R_{ALD} and NO/R_{ALD} , reflecting a broader dependence of χ_{PANs} on NO_x
to aldehydes ratios. For comparison with $\chi_{PANs-obs}$, we include estimates from kinetic calculations assuming a constant $[OH] =$
 2×10^6 molecules cm^{-3} ($\chi_{PANs-kc}$), from calculations using modeled $[OH]$ ($\chi_{PANs-kc w/OH_{model}}$), and from full model
simulations ($\chi_{PANs-model}$) in Fig. 9a. As expected from the kinetic equations, all estimates exhibit a decreasing trend with
580 increasing R_{NO_2}/R_{ALD} , in reasonable accordance with the observations.

Because χ_{PANs} represent a ratio of production rates, it is largely independent of OH concentrations, as indicated by
the overlap between $\chi_{PANs-kc}$ and $\chi_{PANs-kc w/OH_{model}}$ in Fig. 9a. This suggests that the consistently higher $\chi_{PANs-model}$
values than both $\chi_{PANs-kc}$ and $\chi_{PANs-kc w/OH_{model}}$ are not driven by differences in OH but instead by other factors such as
additional PANs sources in the model not accounted for by the kinetic calculations (detailed discussion in Sect. 3.2.1). Despite
585 the similar dependence of observed and estimated χ_{PANs} on R_{NO_2}/R_{ALD} , the estimated values systematically underestimated
 $\chi_{PANs-obs}$. This discrepancy likely reflects the cumulative nature of the observations, which integrate ambient processes such
as total nitrate loss and unaccounted PANs contributions from chemical production, transport, and background sources.

The diagnostic ratio R_{NO_2}/R_{ALD} effectively characterizes the observed partitioning of NO_x loss. To further assess its
utility, we examine the sensitivity of Ln/Q and OPE to changes in R_{NO_2}/R_{ALD} . As shown in Fig. 9b, Ln/Q from model
590 simulations increase with R_{NO_2}/R_{ALD} , indicating shift toward VOC-limited conditions at higher NO_x to aldehyde ratios.
Consistent with this, OPE decreases with increasing R_{NO_2}/R_{ALD} in both kinetic calculations (using $[OH] = 2 \times 10^6$ molecules
 cm^{-3}) and model simulations. These sensitivities likely represent that lower R_{NO_2}/R_{ALD} facilitates radical propagation and PANs
formation, while higher values favor HNO_3 formation and thus radical termination. Although not explicitly stated in kinetic
expressions, R_{NO_2}/R_{ALD} likely evolves with photochemical aging as NO_x is converted to HNO_3 and PANs, and secondary
595 aldehydes are produced downwind. Consistent with this interpretation, OH exposure generally decreases with R_{NO_2}/R_{ALD} (not
shown).

Overall, the dependence of $\chi_{PANs-obs}$ on R_{NO_2}/R_{ALD} , consistent with kinetic theory, suggests continued NO_x
reduction, following the trends observed over the past decade over South Korea and other East Asian countries (Duncan et al.,
2016), may decrease R_{NO_2}/R_{ALD} and shift NO_x loss pathways more toward PANs formation than HNO_3 in winter. Supporting
600 this postulation, $\chi_{PANs-model}$ increased by 22% on average in sensitivity simulations with a 40% reduction in NO_x constraints.
However, this value likely represents an upper limit, as observed aldehydes were constrained in the simulations, despite their
yields from VOC oxidation often being NO_x -dependent (e.g., Millet et al., 2009). The potential shift in NO_x loss partitioning
in response to NO_x reductions may result in extended spatial impacts of NO_x emissions and consequently ozone pollution,
which may be further exacerbated by increase in OPE (Figs. 8 and 9).

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5 Conclusion

This study provides a comprehensive characterization of wintertime acyl peroxy nitrates (PANs) chemistry over South Korea during the Airborne and Satellite Investigation of Asian Air Quality (ASIA-AQ) campaign, combining novel measurements of rare acetyl peroxy nitrates (PAN) homologues with integrated kinetic and modeling analysis. Our findings reveal active photochemical processing despite cold conditions and provide critical insights into volatile organic compounds (VOC) sources, oxidation pathways, and their implications for air quality management.

PANs levels were persistently elevated across South Korea and the Yellow Sea (YS) (frequently exceeding 1 ppbv, maximum 5.5 ppbv), with strong correlations to formaldehyde (CH_2O), odd oxygen (O_x ; approximated as $\text{O}_3 + \text{NO}_2$) and secondary aerosols confirming active winter photochemistry. Regional pollution was extensive, with higher median PANs in remote areas (the mid- and southern region (MS): 990 pptv; YS: 1200 pptv) than in the Seoul Metropolitan Area (SMA: 840 pptv) despite lower emissions, demonstrating significant transport and regional impacts. Elevated homologue/PAN ratios, particularly for acryloyl peroxy nitrates (APAN) and benzoyl peroxy nitrates (PBzN), indicate complex source influences. Notably, strong correlations between APAN, acrolein, and ethylene oxide provided strong evidence of petrochemical impacts and associated toxic exposures in the MS, with back trajectories indicating transport from west coast industrial complexes.

Mechanistic analysis demonstrated that while acetaldehyde dominates PAN production (53-80%), precursors including aromatics make significant contributions (up to 47% in Seoul). Ethanol was demonstrated to be a critical precursor, as it contributes approximately 50% to acetaldehyde formation. In addition, ethanol was very strongly correlated with industrial halocarbons ($r^2 > 0.8$), including bromodichloromethane, HCFC-142b and HFC-134a, indicating important industrial and solvent sources beyond consumer products. This finding challenges previous attributions to volatile chemical products alone and highlights the need for better constraints on industrial VOC emissions in South Korea.

Ozone production analysis confirmed VOC-limited conditions across South Korea, with aldehydes (formaldehyde and C_{2+} aldehydes) contributing approximately 20–30% of instantaneous ozone production. Radical loss was dominated (> 80%) by formation of nitric acid and PANs, with median Ln/Q ratios of 0.9 indicating strong VOC limitation. Ozone production efficiency values (< 10) were consistent with wintertime observations elsewhere in East Asia.

A finding with important policy implications is that continued NO_x reductions may paradoxically expand the spatial extent of ozone pollution. Our diagnostic analysis shows that the fractional contribution of PANs to NO_x loss increases systematically as the ratio of NO_2 to aldehyde reactivity decreases. Model simulations indicate that a 40% NO_x reduction could increase PANs' contribution to NO_x loss by 22%, shifting nitrogen oxide chemistry toward the longer-lived PANs reservoir rather than permanent removal as nitric acid. This shift, combined with increases in ozone production efficiency, may extend the spatial impacts of NO_x emissions and exacerbate regional ozone pollution.

These findings have implications for air quality management in South Korea and East Asia. Effective control of wintertime photochemical pollution requires prioritizing reductions in aldehydes and their precursors—particularly ethanol and reactive alkenes from both industrial and consumer sources. Better characterization of industrial VOC emissions, including from petrochemical facilities, is essential. While concurrent NO_x reductions remain important for long-term benefits and



640 reducing fine particulate matter, policymakers should anticipate potential near-term increases in the spatial extent of ozone
pollution as NO_x levels decline. Finally, our integrated approach demonstrates the value of comprehensive PAN homologue
measurements as diagnostic tools for understanding complex urban and regional photochemistry.

Code and data availability

645 All data used in this work is available at <https://www-air.larc.nasa.gov/cgi-bin/ArcView/asiaaq>. The F0AM model
(version 4.3.0.1) and setup file are available at <https://doi.org/10.5281/zenodo.10069985> (Wolfe et al., 2016).

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Author Contributions

650 All of the authors reviewed and edited the document and participated in the investigation by data acquisition or flight
planning. YRL conceived and performed the formal analysis and wrote the first draft of the document. LA was responsible for
data curation. LGH was responsible for obtaining funding and aided in writing the first draft.

Competing Interests

The authors declare that they have no conflict of interest.

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