

Heat and continental transport shape the variability of volatile organic compounds in the Eastern Mediterranean: Insights from multi-year observations and regional modeling

Anchal Garg¹, Maximilien Desservettaz^{1,2}, Aliko Christodoulou^{1,3}, Theodoros Christoudias¹,
Vijay Punjaji Kanawade¹, Chrysanthos Savvides⁴, Mihalis Vrekoussis^{1,5}, Shahid Naqui¹, Tuija
Jokinen¹, Joseph Byron⁶, Jonathan Williams^{1,6}, Nikos Mihalopoulos^{1,7}, Eleni Liakakou⁷, Jean
Sciare¹, and Efstratios Bourtsoukidis¹

¹Climate and Atmosphere Research Centre, The Cyprus Institute, Nicosia 2121, Cyprus

²Centre for Atmospheric Chemistry, University of Wollongong, Wollongong, NSW 2500, Australia

³PSI, Center for Energy and Environmental Sciences, 5232 Villigen, Paul Scherrer Institute, Switzerland

⁴Department of Labour Inspection, Ministry of Labour and Social Insurance, Nicosia, Cyprus

⁵Institute of Environmental Physics, University of Bremen, Bremen, Germany

⁶Atmospheric Chemistry Department, Max Planck Institute for Chemistry, 55128, Mainz, Germany

⁷Institute for Environmental Research and Sustainable Development, National Observatory of Athens, Palaia
Penteli, 15236 Athens, Greece

*Correspondence: Anchal Garg (a.garg@cyi.ac.cy), Efstratios Bourtsoukidis (e.bourtsoukidis@cyi.ac.cy)

Abstract. Volatile organic compounds (VOCs) are key precursors of tropospheric ozone and secondary organic aerosol formation, yet multi-year observations in the Eastern Mediterranean and Middle East (EMME) remain limited. This study presents multi-year (April 2022-June 2024) high-resolution measurements of 76 VOCs using PTR-ToF-MS at a rural background site in Cyprus, combined with HYSPLIT air-mass analysis to examine the effects of regional transport on VOCs variability. Oxygenated VOCs (OVOCs) dominated the VOCs burden (~79%), followed by aliphatic hydrocarbons, aromatic hydrocarbons, and terpenes. Most VOCs exhibited clear diurnal patterns, observed highest during 08:00-14:00 UTC, varying by species, due to enhanced photochemical activity and temperature-driven emissions. Terpenes, particularly isoprene, increased exponentially with temperature upto 35-38 °C but decreased beyond this threshold, indicating heat-stress inhibition. Monoterpenes showed elevated levels both day and night, reflecting contributions from both biogenic and anthropogenic sources. OVOCs, including acetone, acetaldehyde, methanol, and acetic acid, showed sharp enhancement above 35 °C, consistent with intensified primary emissions and secondary formation under extreme heat. Aromatic hydrocarbons were mainly higher during winter, linked to combustion processes, but benzene levels were highest during summer particularly when temperature rose above 35 °C from evaporative and potential stress-related biogenic sources. HYSPLIT air-mass trajectory analysis revealed dominant contributions from Europe and Northwest Asia (~68%), transporting aged OVOCs, while Middle East winter inflows enhanced aromatic hydrocarbons. While WRF-Chem captured seasonal trends, most VOCs were underestimated, highlighting under-representation of emission sources and oxidation pathways in the model. Overall, the study emphasizes temperature and regional transport as key drivers of VOC variability in the Eastern Mediterranean.

1 Introduction

Volatile Organic Compounds (VOCs) are a diverse group of carbon-based chemicals with high vapor pressures that play a central role in atmospheric chemistry (Atkinson, 2000; Kamal et al., 2016). Upon release into the atmosphere, VOCs participate in photochemical reactions that contribute to the formation of tropospheric ozone (O₃), aerosol precursors, and secondary organic aerosol (SOA), all of which have far-reaching implications for air quality, human health, and climate (Atkinson, 2000; Ehn et al., 2014; Koppmann, 2020; Mellouki et al., 2015; Pennington et al., 2021; Wang et al., 2024). In scientific literature, VOCs are often categorized either by their chemical composition or by their source (Debevec et al., 2017; Koppmann, 2020; Yuan et al., 2024). These compounds with lifetimes ranging from minutes to months

originate from both anthropogenic sources such as fossil fuel combustion, solvent use, industrial processes, and biogenic sources such as vegetation, soils, and marine emissions, and also emissions from biomass burning and wildfires (Atkinson, 2000; Bourtsoukidis et al., 2018; Capes et al., 2009; Civan et al., 2015; Debevec et al., 2017; Desservettaz et al., 2023; Huang et al., 2020; Kaltsonoudis et al., 2016; Pinthong et al., 2022; Pugliese et al., 2023; Ralf Koppmann, 2007; Sindelarova et al., 2014; Wang et al., 2022a; Yuan et al., 2024).

Terpenes (isoprene, monoterpenes, and sesquiterpenes) are primarily emitted (90-95%) by vegetation through enzyme-driven processes influenced by environmental drivers (Bourtsoukidis et al., 2024, 2025; Brito et al., 2018; Ciccioli et al., 2023; Sindelarova et al., 2014; Wang et al., 2024), but some studies (Bryant et al., 2022; Panopoulou et al., 2020) shows that they also have smaller anthropogenic contributions. In the Mediterranean, pine and oak forests are major sources of terpenes (Mecca et al., 2024; Song et al., 2011). Others VOCs such as alkanes, alkenes, and aromatic hydrocarbons, whose major emission sources are anthropogenic arise primarily from fossil fuel combustion, industrial activities, solvent use, biomass burning (both through direct emissions and secondary production) and vehicular traffic, with elevated levels observed in urban areas and near industrial and shipping activities (Atkinson, 2000; Civan et al., 2015; Kajos et al., 2015; Kaltsonoudis et al., 2016; Perrone et al., 2014; Sicre et al., 1987). Oxygenated VOCs (OVOCs) including alcohols, aldehydes, ketones, and organic acids constitute a chemical class with both biogenic and anthropogenic origins, but they are also predominantly formed secondarily via the oxidation of primary VOCs by hydroxyl radicals (OH), O₃, and nitrate radicals (NO₃) (Huang et al., 2020; Mellouki et al., 2015; Wang et al., 2022b). These compounds are important intermediates in the photochemical formation of SOA and contribute to the oxidative capacity of the atmosphere over both local and regional scales (Huang et al., 2020; Liu et al., 2009; Mellouki et al., 2015; Wang et al., 2020, 2022b).

The eastern mediterranean and middle east (EMME) region is a climatically sensitive and chemically dynamic region (Lazoglou et al., 2024; Zittis et al., 2022) owing to high solar radiation, elevated temperatures, and complex meteorology. This region experiences mixed air masses from Western and Eastern Europe, the Middle East, and North Africa, bringing emissions from industrial, biogenic, marine, and biomass burning sources (Bourtsoukidis et al., 2018; Germain-Piaulenne et al., 2024; Lelieveld et al., 2002; Traub et al., 2003). This convergence, combined with the increasing frequency of heatwaves (Lazoglou et al., 2024), enhances VOCs emissions and hence, photochemical activity, raising concerns over regional ozone and SOA formation. Cyprus, located at the core of this region, offers a unique setting to investigate these interactions.

Multi-year measurements of VOCs in the EMME are scarce. At Finokalia, Greece, the studies of Liakakou et al. (2007, 2009) reported extended records of isoprene and C₂-C₈ hydrocarbons, while in Athens, the studies of Panopoulou et al. (2018, 2020) presented multi-year observations of isoprene, terpenes, and C₂-C₈ non-methane hydrocarbons (NMHCs). In contrast to these gas chromatography-based observations, proton-transfer-reaction time-of-flight mass spectrometry (PTR-ToF-MS) studies in the region have thus far been limited to short-term, campaign-based deployments (Derstroff et al., 2017; Kaltsonoudis et al., 2016). Overall, despite extensive research in diverse regions, the understanding of VOCs variability and drivers in

the EMME region remains limited, particularly in terms of long-term and multi-year observations that can capture seasonal and climate change-driven dynamical responses to extreme heat.

Addressing this gap, the current study presents multi-year high-resolution VOCs measurements collected from 2022 to 2024 at a rural background site in Cyprus located in EMME region. Using PTR-ToF-MS, we quantified and classified 76 VOCs. To identify regional transport, we apply hybrid single-particle lagrangian integrated trajectory (HYSPLIT) air mass back-trajectory analysis and examine air mass origins across seven distinct geographical sectors. Furthermore, we evaluated selected VOCs simulated by the weather research forecast model coupled with chemistry (WRF-Chem) model which includes a high-resolution mapping of local emissions (Georgiou et al., 2022). This integrated approach enables us to (i) characterize diurnal and seasonal VOCs patterns, (ii) assess the influence of temperature on VOCs composition, (iii) understand key emission sources, (iv) evaluate the impact of long-range transport on VOCs variability, and (v) assess the model performance in the estimation of VOCs.

2 Methodology

2.1 Study area and VOCs measurement timeline

Cyprus, the third-largest island in the Mediterranean is situated at the air pollution crossroads of Europe, Asia, and Africa, making it a receptor for long-range atmospheric transport from diverse source regions (Lelieveld et al., 2002). Air masses influencing the island originate from Western and Eastern Europe, Turkey, North Africa, Southwest and Northwest Asia, the Middle East, and the surrounding Mediterranean Sea (Fig. 1a). This convergence of continental and marine influences makes Cyprus an ideal location for investigating the chemical complexity of both long-range transported and local air pollutants (Meusel et al., 2016; Derstroff et al., 2017; Debevec et al., 2017; Vrekoussis et al., 2022).

The present study was conducted at the Cyprus Atmospheric Observatory-Agia Marina Xyliatou (CAO-AMX, 35.0387° N, 33.0579° E), located at 532 m above mean sea level at the foothills of the Troodos Mountains (Fig. 1b-c). Established in 1997 and upgraded in 2015 to a “supersite,” CAO-AMX provides high-quality, long-term measurements of aerosols and their properties, greenhouse gases, meteorological parameters, and trace gases. It is designed and operated as a rural background site (CAO-AMX, <https://cao.cyi.ac.cy/>) primarily by the climate and atmosphere research center (CARE-C) of the Cyprus institute (CyI), in collaboration with national authorities such as the department of labour inspection (DLI) of the ministry of labour and social insurance, while it is part of major international atmospheric monitoring networks, including ACTRIS (<https://www.actris.eu/>); EMEP (<https://www.emep.int/>); and AERONET (<https://aeronet.gsfc.nasa.gov/>) (Sciare J, 2016). The station is situated near the small villages of Agia Marina (~630 inhabitants) and Xyliatos (~150 inhabitants), surrounded by oak-pine forests and Maquis shrubland, with agricultural fields ~4 km to the north (Baalbaki et al., 2021). The nearest urban area lies >35 km away, ensuring minimal influence from local pollution sources aside from limited vehicle traffic. Its rural setting, surrounded by forested and low-population areas and situated far from major urban sources (Nicosia about 35 km to the northeast and Larnaca about 50 km to the southeast) ensures that measurements are minimally influenced by major nearby local anthropogenic emissions. This allows the observation of representative background concentrations and the signatures of regional and long-range transported pollutants.

VOCs observations were conducted at the CAO-AMX site from 27 April 2022 to 15 June 2024, with occasional interruptions caused by other field campaigns or instrument downtime. Meteorological data were recorded at a temporal resolution of 5 minutes from a dedicated station located in the village of Xyliatos, approximately 2.85 km from the primary observational site at the CAO-AMX. DLI managed the installation and operation of a relatively dense monitoring network at this station. As shown in Fig. 1d, monthly data coverage varied but provided efficient coverage across all seasons. Fig. 1e summarizes the seasonal and annual distribution of data points: summer (32%) was the most represented season, followed by spring (30%) and autumn (23%), while winter accounted for 15% of the data. In terms of annual contribution, 2022 comprised 44% of the total dataset, 2023 contributed 25%, and 2024 contributed 31%.

2.2 VOCs measurements

Online measurements of VOCs were performed using a high-resolution PTR-ToF-MS (Model: 4000; Ionicon Analytik GmbH, Innsbruck, Austria) following the methodology as described by Desservettaz et al., (2023). PTR-ToF-MS is a high-sensitivity, real-time technique that measures VOCs by transferring a proton from H_3O^+ ions to VOC molecules. It provides rapid, accurate, and high-resolution detection of a wide range of atmospheric VOCs without sample preparation.

Briefly about the methodology for VOCs measurement, the drift tube of the PTR-ToF-MS was operated under controlled conditions with an electric field to number density ratio (E/N) of 128 Townsend (Td), a pressure of 2.2 mbar, an applied voltage of 630 V, and a temperature of 85 °C. Ambient air was sampled at a flow rate of 210 mL min⁻¹ through a 2.0 m long, 1/16" PEEK (polyether ether ketone) heated inlet line, equipped with a 0.45 µm PTFE particulate filter (PALL, USA). The inlet drew air from a common glass manifold shared with other atmospheric measurement instruments. The use of chemically inert materials such as PTFE and PEEK minimized potential compound losses within the sampling line.

The instrument was automatically switched to measure VOC-free air for 20 minutes after every 23 hours, followed by 40 minutes of calibration using a certified gas standard (Apel-Riemer Environmental, USA) containing major VOCs diluted to ~20 ppbv; this 23-h cycle caused the non-measurement period to shift by approximately 1 h each day, ensuring that no specific time of day was systematically underrepresented and that diurnal patterns were not biased. VOCs not included in the gas standard were calibrated using established relative response factors determined from prior laboratory experiments and, where possible, verified with secondary standards. Mass spectra were acquired at 1-minute time resolution and averaged to 1-hour intervals for the analysis in this study. The instrument sensitivity was interpolated between calibrations and adjusted for humidity dependence based on water cluster ion ratios. Calibrated sensitivities showed modest temporal variability (~10-23%) for most compounds. Post-acquisition analysis was carried out using the Ionicon data analyzer (IDA), with mass axis calibration based on H_3O^+ and internal reference peaks (e.g., $\text{C}_6\text{H}_5\text{I}^+$). The software retrieved exact masses and chemical formulas for ~1200 peaks, of which 400–500 was assigned tentative structures using the GLOVOCS database, which is a master compound assignment guide that can be referred to as PTR-MS practitioners (Yáñez-Serrano et al., 2021). A set of 131 common species across analysis periods was further corrected using transmission curves derived from zero and calibration air sampling. Mixing ratios (ppbv) were calculated from ion counts, incorporating species-specific reaction rate constants (k-values) based on molecular dipole moments and polarizabilities

(Desservettaz et al., 2023; Pagonis et al., 2019). The limit of detection (LOD) for each species was defined as three times the standard deviation observed during zero-air measurements and averaged over the entire campaign. VOC species with more than 30% of their data below the LOD were excluded, resulting in a final dataset of 76 species, with sub-LOD concentrations appropriately flagged. Measurements impacted by filter changes, exhibiting short-term interference (notably acids and certain hydrocarbons), were filtered by removing a conservative 2-hour window following each swap. Formaldehyde was not included in this study due to its inconsistent identification by the IDA software during part of the measurement period. Furthermore, quantification of formaldehyde via PTR-based techniques is known to be highly sensitive to humidity variations (Vlasenko et al., 2010), which complicates reliable detection. Accurate quantification would require water-dependent calibrations and a dedicated formaldehyde calibration standard, conditions that were not met during the measurement period.

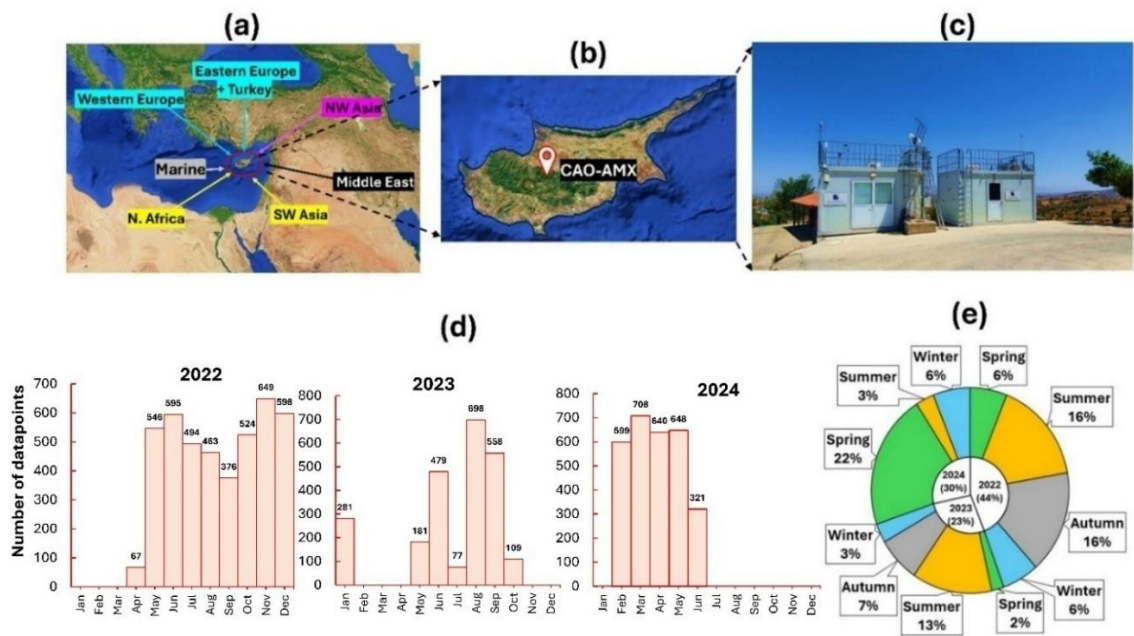


Figure 1: Geographic context, air mass origins, and VOC measurement coverage at the CAO-AMX site (2022-2024). (a) Major regional air mass source areas influencing Cyprus; (b) Location of the CAO-AMX in the Troodos Mountains; (c) The CAO-AMX monitoring station, a designated regional background site; (d) Monthly distribution of number of datapoints for VOC measurements from April 2022 to June 2024; (e) Summary of data coverage by season and year, showing overall representativeness of the dataset for multi-seasonal and multi-annual analysis.

2.3 Meteorological and air pollutant data

Meteorological and auxiliary air quality parameters were analyzed to provide contextual insight into atmospheric dynamics and pollutant characteristics. The meteorological station (mentioned in section 2.1) continuously measured key meteorological parameters including ambient temperature (AT), relative humidity (RH), solar radiation (SR), wind speed (WS), and wind direction (WD). In addition, a range of trace gases and particulate pollutant levels were measured at a co-located air quality monitoring station (~20 m from CAO-AMX, main VOCs measurement container), operating under the European monitoring and evaluation programme (EMEP) framework provides hourly-resolved concentrations of major atmospheric pollutants: ozone (O_3), carbon monoxide (CO), nitrogen oxides (NO , NO_2 , and NO_x), sulfur dioxide (SO_2), and particulate matter with aerodynamic diameters less than $10 \mu m$ (PM_{10}) and $2.5 \mu m$ ($PM_{2.5}$). These

measurements serve as critical auxiliary data for assessing the chemical background of the region, evaluating pollutant transport processes, and validating model simulations. Instrument specifications, calibration protocols, and operational details of the meteorological and air quality monitoring systems are extensively documented in previous studies (Kleanthous et al., 2014; Pikridas et al., 2018; Vrekoussis et al., 2022).

2.4 HYSPLIT backward trajectory

To assess the influence of long-range atmospheric transport on local air quality, five-day back trajectories were computed using the HYSPLIT model (version 4.4; Draxler, 1997; Stein et al., 2015). These calculations were performed using the PC-based version of HYSPLIT provided by the national oceanic and atmospheric administration's (NOAA) air resources laboratory (ARL), incorporating meteorological data from the global data assimilation system (GDAS) with a spatial resolution of $1^\circ \times 1^\circ$ and a temporal resolution of 1-hour (Draxler, 1997). Trajectories were initialized every hour at an altitude of 500 m above ground level (AGL), and each point along the trajectory was calculated at a one-hour time step. The air mass trajectory cluster analysis, following the methodology previously applied by Debevec et al., (2017) identified a total of seven air mass clusters: (C0) slow-moving or stagnant local air masses, (C1) North Africa, (C2) Marine-influenced, (C3) Europe, (C4) Northwest Asia, (C5) West Turkey, and (C6) Middle east origins. For the cluster analysis, the altitude of the end point was set at 1000 m AGL for each trajectory (Debevec et al., 2017).

2.5 WRF-Chem model configuration for VOC simulation in Cyprus

To assess VOCs variability and compare modeled and observed mixing ratios over Cyprus, we employed the WRF-Chem model as described and evaluated in Georgiou et al. (2022). The setup used three nested domains with horizontal resolutions of 50, 10, and 2 km, the later focusing on Cyprus, with meteorological inputs from the global forecast system (GFS) every 3 hours. Key parameterizations included Yonsei University (YSU) planetary boundary layer (PBL) scheme for boundary layer, Noah land surface model (LSM) for land surface processes, the rapid radiative transfer model for global applications (RRTMG) for radiation, the regional atmospheric chemistry mechanism (RACM) for gas-phase chemistry and Modal Aerosol Dynamics for Europe (MADE) for inorganic aerosols and Volatility Basis Set (VBS) for organic aerosols as described in Georgiou et al. (2022). Photolysis was calculated using Fast-J. Anthropogenic emissions were sourced from emissions database for global atmospheric research-hemispheric transport of air pollution (EDGAR-HTAP; version 2) emission inventory, while BVOCs were simulated online using the Model of Emissions of Gases and Aerosols from Nature version (MEGAN) version 2.1 based on weather and land use data (Guenther et al., 2012).

For the innermost domain, a high-resolution emission inventory developed by Georgiou et al. (2020) is used. This emission inventory uses the total reported emissions of CO, NO_x, non-methane VOCs (NMVOCs), SO₂, and PM on a $1 \text{ km} \times 1 \text{ km}$ resolution (Georgiou et al., 2020) which is upscaled to the resolution of the innermost domain of the simulations (2 km) using a nearest-neighbor grid-point attribution algorithm, while diurnal, weekly, and monthly emission cycles are applied to each species and the predominant emission activity per season according to the (Schaap et al., 2011). The high -resolution emission inventory was found to adequately simulate of the daily profiles of NO_x and O₃ at measurement sites over Cyprus, and compared

to CAMS, the WRF-Chem model predicts more accurately NO₂ mixing ratios with 7% during winter and -44 % during summer normalized mean bias during winter/summer, compared to -81% and -84% biases for the CAMS ensemble using the CAMS inventory (Georgiou et al., 2022). In the outer model domain, the EDGAR emission inventory is used for anthropogenic emissions, and MEGAN for biogenic emissions. In comparison with other emission inventories for VOCs, EDGAR provides a larger estimate than the EMEP emission inventory. The CAMS-REG-AP inventory also gives a smaller estimate, in comparison to EMEP. Comparing the VOC by the EDGAR inventory to the EMEP inventory, the higher emission estimates are greatest over Belgium, Austria, Switzerland, Germany and Finland, with differences mainly encountered in the industrial combustion sector and fugitive emissions. Regarding the modelled O₃ concentrations over Europe, all emission inventories were shown very similar patterns despite the noted differences in terms of emissions, particularly VOCs, indicating a low sensitivity to VOCs emissions (Thunis et al., 2021).

Major VOCs such as isoprene, monoterpenes, and aromatics were explicitly modeled. Outputs at hourly resolution were validated against PTR-ToF-MS measurements from CAO-AMX, enabling robust interpretation of VOC sources and atmospheric transport in the eastern mediterranean. The VOCs included in this study are based on the RACM (Stockwell et al., 1997) and encompass key anthropogenic and biogenic compounds relevant to regional air quality. These include isoprene (ISO), monoterpenes represented by α -pinene and related monocyclic terpenes with a single double bond (API), and d-limonene, along with other cyclic dienes (LIM). Aromatic compounds are grouped as toluene (TOL), which includes toluene and less reactive aromatics, and xylenes (XYL), representing xylenes and more reactive aromatic species. OVOCs are categorized as aldehydes (ALD), ketones (KET), and glyoxal (GLY).

3 Results and Discussion

3.1 Meteorological and atmospheric composition variability

Mean meteorological conditions included WS of $2.6 \pm 1.4 \text{ m s}^{-1}$, RH of $55 \pm 20\%$, AT of $17.7 \pm 7.9 \text{ }^{\circ}\text{C}$, and SR of $224.75 \pm 305.85 \text{ W m}^{-2}$ (Fig. 2a). Average AT ($>30 \text{ }^{\circ}\text{C}$) and SR ($>800 \text{ W m}^{-2}$) peaked during daytime in summer, while winter had the lowest daytime values ($<15 \text{ }^{\circ}\text{C}$ and $<400 \text{ W m}^{-2}$ Fig. 2e-f). RH showed the opposite pattern during daytime ($\sim 60\%$ in winter, $<40\%$ in summer) (Fig. 2g). Midday peaks in temperature and radiation were strongest in summer, with RH lowest in the afternoon (Fig. 2e-g). RH and SR were inversely correlated, consistent with previous studies (Emekwuru and Ejohwomu, 2023; Matthew, 2022). Daily average WS ranged from 1 to 4 ms^{-1} , with slightly higher values observed in spring and summer, suggesting enhanced mixing during these periods (Fig. 2h). Diurnal patterns were also pronounced as WS showed afternoon maxima in all seasons, with enhanced values in spring and summer, likely driven by thermal convection and sea breeze circulation (Fig. 2h), promoting more effective dispersion of air pollutants (Deot et al., 2025; Pérez et al., 2020; Soukissian and Sotiriou, 2022).

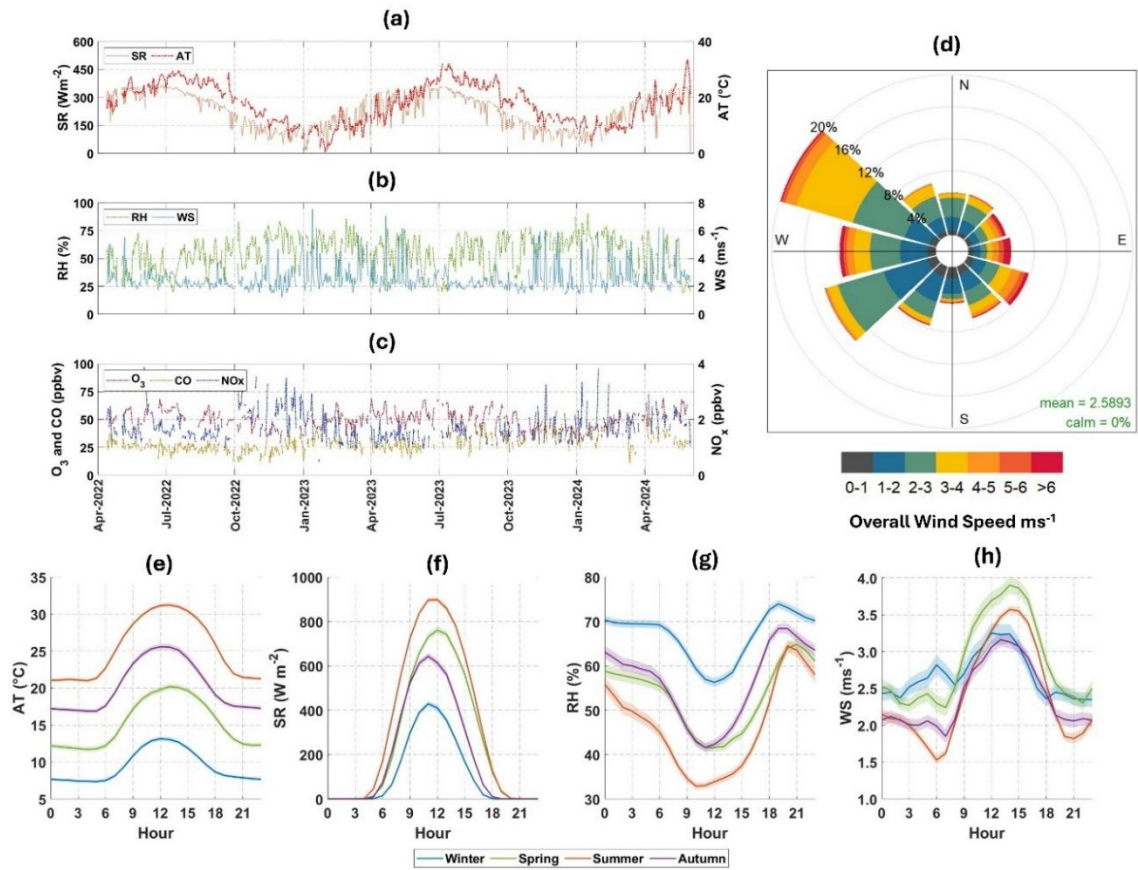


Figure 2: Seasonal and diurnal variability of meteorological parameters and air pollutants in Cyprus. Panels (a-c) show time series (24-hour rolling average) of (a) ambient temperature (AT) and solar radiation (SR), (b) relative humidity (RH) and wind speed (WS), and (c) gaseous pollutants, including ozone (O_3), carbon monoxide (CO), and nitrogen dioxide (NO_2) for study period. Panel (d) display wind rose diagrams indicating wind speed and direction for the entire study period. Panels (e-h) present the diurnal profiles of (e) ambient temperature ($^{\circ}C$), (f) solar radiation ($W m^{-2}$), (g) relative humidity (%), and (h) wind speed (ms^{-1}) across the four seasons (spring, summer, autumn, winter). Shaded regions in panels (e-h) represent the standard error, capturing variability in diurnal cycles.

Wind analysis throughout the study period revealed predominantly moderate conditions, with over 70% wind speeds falling below $3 ms^{-1}$, and 94% below $5 ms^{-1}$. The wind rose diagrams (Fig. 2d) highlight the prevalence of moderate-speed winds and dominant directions ranging from northwesterly to west-southwesterly. Fig. S1 shows seasonal and monthly variability in wind direction, and suggests that during the winter, the winds were mainly coming from the southeast, followed by the southwest, with most wind speeds exceeding $4 ms^{-1}$; whereas, for the other seasons, the winds originated predominantly from the northwest.

Atmospheric conditions were characterized by the following mean mixing ratios for gaseous pollutants such as NO at 0.3 ± 0.2 ppbv, NO_2 at 1.4 ± 0.8 ppbv, NO_x at 1.7 ± 0.9 ppbv, SO_2 at 0.5 ± 0.4 ppbv, O_3 at 49.2 ± 9.0 ppbv, and CO at 145.8 ± 49.5 ppbv. During daytime O_3 remained consistently elevated throughout the year, with a significant increase of (8-20 ppbv) of its mixing ratio in summer, indicating a strong influence from regional transport and photochemical formation (Kleanthous et al., 2014; Lee et al., 2021; Pochanart et al., 2003). A notable increase in CO levels was observed between 2022 and 2024, particularly linked to increase in anthropogenic activities. A modest rise in CO observed during winter is likely due to increased combustion emissions, lower sink due to low photochemistry, variable residence time of upto 90 days in winter and 30 days in summer, and weaker atmospheric dispersion (Koppmann et al., 2005; Pochanart et al.,

2003). NO_x showed a winter maximum, consistent with increased anthropogenic emissions and reduced photochemical degradation (Atkinson, 2000; Civan et al., 2015; Koppmann et al., 2005).

3.2 Overview of atmospheric VOCs observations

285 A total of 76 VOCs were quantified, and their descriptive statistics for key species (the most studied VOCs
in EMME region and European studies) are summarized in Table 1, while the complete list is provided in
supplementary Table S1. These VOCs were grouped into five major chemical classes: (i) OVOCs (alcohols,
aldehydes, ketones, acids, and other oxygenates); (ii) aliphatic hydrocarbons; (iii) aromatic hydrocarbons;
290 (iv) terpenes; and (v) nitrogen- and sulfur-containing VOCs. An overview of the average mixing ratios is
shown in Fig. 3, while Fig. S2 illustrates the time series of selected VOCs. The selected VOCs shown in
Table 1 were most studied in the previous Mediterranean and European studies, whereas Fig S3 shows the
20 most abundant VOCs observed in this study.

OVOCs were the dominant chemical class as also observed by Debevec et al. (2017). In our study OVOCs
contributed approximately 79% to the total measured VOC burden. Among these, alcohol was the most
295 abundant subgroup, comprising 32% of total VOCs. Methanol was found as the single most dominant
species, with a median mixing ratio of 2.86 ppbv, accounting for 87% of the total alcohol fraction. Its high
dominance reflects oxidation of biogenic and biomass burning emissions, consistent with previous findings
(Debevec et al., 2017; Mukherjee et al., 2024; Yuan et al., 2024). Carbonyls (aldehydes, and ketones)
represented the second largest OVOCs subgroup (28% of total VOCs), with median levels dominated by
300 acetone (2.11 ppbv, 74% of the carbonyl group), followed by acetaldehyde (0.54 ppbv, 19% of the carbonyl
group). These compounds are known to originate from both primary sources and secondary oxidation of
VOCs (Capes et al., 2009; Mellouki et al., 2015; Mukherjee et al., 2024; Pennington et al., 2021; Wang et
al., 2024, 2022b). Organic acids, dominated by acetic acid (median: 0.68 ppbv) and formic acid (median:
0.42 ppbv), contributed 13% of total VOCs. Their contributions likely arise from a combination of direct
305 biogenic and anthropogenic emissions, as well as secondary formation, which could not be quantitatively
distinguished. (Derstroff et al., 2017; Huang et al., 2020; Kamal et al., 2016; Koppmann, 2020; Wang et al.,
2024, 2022b). Other oxygenates, such as methyl ethyl ketone (MEK) and acrolein, also contributed to the
OVOCs group and may originate from industrial sources and atmospheric oxidation (Schieweck et al., 2021;
Torres-Vinces et al., 2020; Yáñez-Serrano et al., 2016).

310 Aliphatic hydrocarbons accounted for 12% of total VOCs, with species such as propylene, propyne, 1,3-
butadiene, and cyclohexadiene likely reflecting emissions from vehicular traffic, fossil fuel combustion and
industrial activities (Perrone et al., 2014; Sicre et al., 1987). Aromatic hydrocarbons, though less abundant
(3%), were dominated by benzene (0.14 ppbv, 32% of the aromatic fraction), followed by toluene and
xylenes. These compounds are well-known tracers of urban and industrial emissions (Abbasi et al., 2020;
315 Bourtsoukidis et al., 2019; Bourtsoukidis et al., 2020; Das et al., 2024), though their overall concentrations
were low, consistent with the remote and background nature of the site.

Terpenes, such as monoterpenes (median= 0.27 ppbv) and isoprene (median= 0.09 ppbv), collectively
contributing ~3% of the VOC budget. Their mixing ratio peaked in summer, aligning with increased biogenic

emissions driven by higher temperatures and SR (Gu et al., 2021; Liakakou et al., 2007; Mukherjee et al., 2024; Sindelarova et al., 2014; Strada et al., 2023; Wang et al., 2024).

Nitrogen- and sulfur-containing VOCs also contributed approximately 3%, with median acetonitrile (0.11 ppbv) and dimethyl sulfide (DMS, 0.04 ppbv) mixing ratios as the major constituents. Acetonitrile, a tracer for biomass burning, accounted for 40% of this group, while DMS accounted for 17% of this group reflects both marine and biological sources (Deschaseaux et al., 2022; Edtbauer et al., 2020; Huangfu et al., 2021; Robles, 2005).

The analysis of the 20 most abundant VOCs (Fig. S3) also reinforced the dominance of OVOCs, particularly methanol, acetone, and acetic acid, which together formed a substantial fraction of the observed VOC mass. The relatively low levels of aromatics and aliphatic further support the interpretation that this rural background site is primarily influenced by transported VOC precursors, secondary atmospheric processes and regional-scale transport, rather than direct anthropogenic emissions. A major reason for the low burden of aliphatic hydrocarbons in the PTR-ToF-MS measurements is their inherently low proton affinities, which result in weak ionization and significant fragmentation. Consequently, their signals are unreliable or very small, preventing a clear aliphatic hydrocarbon profile from being obtained in this study.

Table 1. Descriptive statistics of selected VOCs (in ppbv) measured in Cyprus (April 2022- June 2024)

Chemical Name	Chemical Formula	Chemical Class	M.H ⁺	Min	Max	Mean	Median	SD
Methanol	CH ₄ O	Alcohol	33.03	0.138	27.82	3.08	2.86	1.71
Acetonitrile	C ₂ H ₃ N	N-containing VOC	42.04	0.041	0.33	0.12	0.11	0.04
Acetaldehyde	C ₂ H ₄ O	Aldehyde	45.03	0.123	6.76	0.88	0.54	0.89
Formic Acid	CH ₂ O ₂	Acids	47.01	0.022	7.99	0.78	0.42	0.93
Ethanol	C ₂ H ₆ O	Alcohol	47.05	0.009	3.68	0.43	0.35	0.35
Acrolein	C ₃ H ₄ O	Oxygenated	57.03	0.01	1.93	0.2	0.12	0.23
Glyoxal	C ₂ H ₂ O ₂	Oxygenated	59.01	0.019	1.22	0.07	0.07	0.04
Acetone	C ₃ H ₆ O	Ketone	59.05	0.429	10.82	2.35	2.11	1.19
Acetic Acid	C ₂ H ₄ O ₂	Acids	61.03	0.039	12.54	1.25	0.68	1.56
DMS	C ₂ H ₆ S	S-containing VOC	63.04	0.005	0.55	0.06	0.04	0.06
Furan	C ₄ H ₄ O	Oxygenated	69.03	0.004	0.16	0.01	0.01	0.01
Isoprene	C ₅ H ₈	Terpenes	69.07	0.005	2	0.15	0.09	0.18
MVK/Methacrolein	C ₄ H ₆ O	Oxygenated	71.05	0.003	0.53	0.08	0.06	0.07
MEK	C ₄ H ₈ O	Oxygenated	73.07	0.007	2.78	0.36	0.29	0.27
Benzene	C ₆ H ₆	Aromatics	79.05	0.018	1.09	0.16	0.14	0.11
Toluene	C ₇ H ₈	Aromatics	93.07	0.005	1.3	0.1	0.07	0.1
Xylenes	C ₈ H ₁₀	Aromatics	107.09	0.003	1.76	0.06	0.03	0.12
Monoterpenes	C ₁₀ H ₁₆	Terpenes	137.13	0.026	3.01	0.34	0.27	0.27

Compounds are grouped by chemical class, with molecular hydronated ion mass (M.H⁺) shown for each. These data provide insight into the variability and typical concentrations of VOCs in the region, relevant to air quality and atmospheric chemistry assessments. These are the most studied VOCs in EMME region and European studies. For clarity, nitrogen and sulfur containing VOCs class has been separated into distinct classes, N-containing VOC and S-containing VOC in this table to explicitly highlight the nitrogen contribution in acetonitrile and the sulfur contribution in DMS.

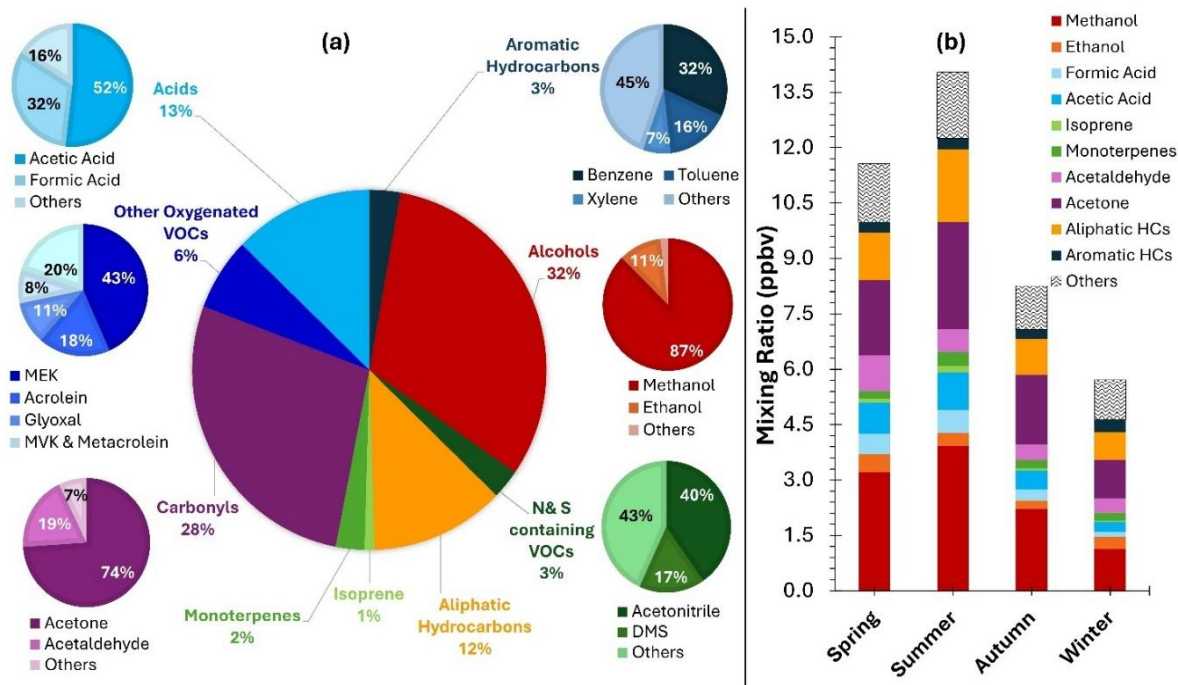


Figure 3: Overview of VOCs in Cyprus: Composition and seasonal variability in its composition. (a) % contribution of different VOC groups in Cyprus, highlighting the dominance of alcohols (32%), carbonyls (28%), and acids (13%). Subcategories within each VOC group are detailed in smaller pie charts. (b) Seasonal variability of the VOCs mixing ratio. Here carbonyls include aldehydes, and ketones. Overall OVOCs include alcohol, carbonyls, acids, and other oxygenated VOCs. Isoprene and Monoterpenes combine together to form terpenes class.

As shown in Table 2, VOC mixing ratios measured in our study are within the range reported from other European background and rural sites. Winter mixing ratios for acetaldehyde, acetone, MVK, MEK, acetonitrile, benzene, isoprene, and monoterpene were also comparable to those measured during the same period in 2015 at this site (Debevec et al., 2017), suggesting stable background concentrations and low interannual variability in the Eastern Mediterranean. However, summertime isoprene mixing ratios at Ineia, Cyprus, were about three times lower. Derstroff et al. (2017) demonstrates potential contribution of downwind forest emissions or the effect of hotter summers and stress induced emissions. Elevated levels of most VOCs during summer (Fig. 3b) suggest the influence of increased biogenic activity and enhanced photochemical production, as well as contributions from solvent use and fuel evaporation.

Table 2. Seasonal mean mixing ratios (ppbv) of VOCs measured in our study compared with previous regional and European studies.

Location	Measuremen t	Study Reference	Year, Season	Terpenes		Aromatic Hydrocarbons			Oxygenated VOCs							N-containing VOC
				Isoprene	Mono- terpenes	Benzene	Toluene	Xylenes	Methanol	Ethanol	Acetic Acid	Acetaldehyde	Acetone	MVK+ Methacrolein	MEK	Acetonitrile
CAO, Cyprus (Background)	PTR-ToF- MS	This study	Spring 2022-24	0.12	0.20	0.14	0.06	0.03	3.22	0.47	0.84	0.97	2.03	0.06	0.35	0.10
			Summer 2022-24	0.18	0.39	1.30	0.07	0.03	3.92	0.35	1.00	0.61	2.90	0.09	0.28	0.13
			Autumn 2022-24	0.06	0.23	0.12	0.06	0.02	2.22	0.23	0.52	0.42	1.90	0.04	0.24	0.12
			Winter 2022-24	0.03	0.22	0.15	0.09	0.04	1.14	0.33	0.27	0.39	1.05	0.03	0.29	0.08
CAO, Cyprus (Background)	PTR-MS, GC-MS	Debevec et al., 2017	Winter, 2015	0.05	0.24	0.12	0.05	0.02	2.93			0.41	1.15	0.03	0.22	0.12
Ineia, Cyprus (Background)	PTR-MS	Derstroff et al., 2017	Summer, 2014	0.06	0.11	0.02	0.01	0.02	2.90		0.95	0.32	2.25	0.03	0.11	0.11
Zurich, Switzerland (Urban background)	GC-MS	Legreid et al., 2007	Spring 2005	0.08		0.41	1.46	0.97	2.18	6.87			1.66	0.04	0.24	
			Summer 2005	0.16		0.23	1.43	0.83	3.18	3.94		0.80	2.12	0.12	0.20	
			Autumn 2005	0.08		0.48	1.70	1.10	1.11	7.61		0.45	1.24	0.04	0.17	
			Winter 2005	0.06		0.75	1.25	0.91	1.21	7.53		0.82	1.17	0.05	0.22	
Athens, Greece (urban)	PTR-MS	Kaltsonoudis et al., 2016	Summer 2012	0.73	0.92	0.22	0.81	0.67		1.52	2.17		4.28	0.35	0.50	0.20
			Winter 2013	1.05	0.43	1.00	2.34	1.69		1.80	2.11		2.24	0.41	0.59	0.16
Cape Corsica, France (urban)	PTR-MS	Michoud et al., 2017	Summer 2013	0.19	0.41	0.03	0.08			0.18			3.43	0.06	0.48	
Oak Mediterranean Forest, France	PTR-MS	Kalogridis et al., 2014	Spring 2012	1.19	0.06	0.07	0.05		2.28			0.38	1.28	0.21		
Oak Mediterranean Forest, Spain	PTR-ToF- MS	Seco et al., 2011	Summer	0.15- 0.75	0.13- 1.42	0.04- 0.09	0.08- 0.47		4.14-6.05	1.15- 2.87	1.32- 2.94	0.54-1.26	2.26- 3.83	0.17-0.54		0.16-0.22
			Winter	0.02- 0.07	0.005- 0.067	0.11- 0.19	0.06- 0.41		1.28-2.70	0.47- 2.08	0.39- 1.43	0.23-0.66	0.79- 1.55	0.01-0.05		0.08-0.10
Barcelona, Spain (urban background)	PTR-ToF- MS	In't Veld et al., 2024	Spring 2022	0.04	0.05	0.14	0.47		3.50			0.92	1.93	0.09	0.22	0.41
Montseny, Spain (rural background)			Summer 2022	0.18	0.19	0.08	0.80		3.73			0.31	2.50	0.17	0.24	0.29
Montseny, Spain (rural background)			Spring 2022	0.05	0.02	0.13	0.22		1.46			0.48	1.02	0.05	0.19	0.10
			Summer 2022	0.30	0.45	0.04	0.09		3.89			0.84	2.76	0.55	0.30	0.12
Oak Mediterranean Forest, Spain	PTR-MS	Yáñez-Serrano et al., 2021	Summer 2019	0.42	0.15	0.04	0.13		4.60			0.77	2.08	0.30	0.29	0.12
			Autumn 2019	0.09	0.09	0.04	0.21		1.60			0.30	1.37	0.08	0.27	0.06

360 As shown in Table 2, the Mediterranean forested sites in France and Spain are characterised by
higher daily mean isoprene levels up to 1 ppbv (Kalogridis et al., 2014; Seco et al., 2011), consistent
with strong direct emissions from oak-dominated ecosystems compared to the conifer-dominated
forest in Cyprus. The urban site of Zurich, Switzerland shows similar seasonal pattern as our study,
365 where isoprene, methanol, acetone, and MVK all showed a summer maximum and a winter
minimum (Legreid et al., 2007). However, wintertime aromatic (BTX) concentrations were much
higher in Zurich presumably due to combustion sources related to anthropogenic activities. Nitriles
and MEK concentrations in our study fall in the range of those reported for previous European
background studies (Table 2) and confirm their regional well-mixing as resulting from both
anthropogenic and biomass-burning sources.

370 3.3. Seasonal and diurnal variability of VOCs functional groups

The diurnal variation of total VOCs (TVOCs) across all seasons and years (Fig. S4) reveals clear
patterns influenced by temperature, photochemistry, boundary layer dynamics, and emission
sources. TVOCs mixing ratio typically observed high between 09:00-14:00 UTC, corresponding to
enhanced daytime emissions and increased atmospheric reactivity. The highest median TVOCs
375 mixing ratios were observed during summer 2024 (25.76 ppbv), followed by summer 2023 (16.27
ppbv), spring 2024 (13.71 ppbv), and summer 2022 (12.35 ppbv), whereas winter exhibited the
lowest levels (5.83 ppbv). The less pronounced seasonal enhancement observed in summer 2022
compared to 2023 and 2024 is likely due to comparatively lower ambient temperature of about 1-3
°C resulting in weaker photochemical activity during that period, and reduced temperature-driven
380 biogenic emissions. In contrast, summer 2024 was characterized by exceptional heat, including
record-breaking temperatures (>45°C) across Cyprus, which strongly intensified direct VOCs
emissions and secondary production.

June 2024 was the warmest June on record in Cyprus, with 43 of 52 stations registering record
maximum temperatures; during the 14–15 June heatwave, maximum temperature reached 45.3 °C,
385 resulting in two heat-related deaths and widespread wildfire impacts (Department of Meteorology,
Cyprus, 2024). During summer, there is also northerlies that promote the transport of species from
regions with pronounced anthropogenic emissions, including biomass burning, contributing to
elevated levels (Vrekoussis et al., 2022). The prolonged heat (summer 2024) was attributed to a
persistent heatwave influenced by global climate change, with such extreme events forecasted to
390 become more frequent and intense in the future (Yáñez-Serrano et al., 2016; Zittis et al., 2022).
Similarly, the higher VOCs mixing ratio measured during spring 2024 relative to earlier spring
campaigns can also be attributed to warmer-than-usual conditions, which enhanced biogenic release
and photochemical reactivity. Fig. 4 presents the diurnal profiles of VOCs for multiple seasons
(2022–2024), which demonstrate pronounced variability on seasonal and diurnal dynamics,
395 particularly during summer 2024, when elevated temperatures significantly enhanced VOCs levels
across most chemical classes.

3.3.1. Terpenes (isoprene and monoterpenes)

High isoprene levels occurred during 08:00-14:00 UTC (the time when SR and AT are highest) in summer 2023 and 2024 (Fig. 4a).. In summer 2024, the median isoprene mixing ratio reached approximately 1.0 ppbv during the day, with daily average median values of 0.43 ± 0.36 ppbv. Interestingly, the observed isoprene maximum at 08:00 UTC does not coincide with the peaks in SR and AT. This suggests that under the extreme drought conditions of 2023 and 2024, vegetation may have downregulated photosynthesis unusually early in the day. Such an early shutdown indicates that a physiological threshold was crossed, whereby plants shift their carbon allocation strategy from growth toward survival. While drought stress is often associated with an initial increase in isoprene emissions (Byron et al., 2022), prolonged or severe stress can suppress photosynthesis and consequently reduce isoprene release (Potosnak et al., 2014; Byron et al., 2022). On the other hand, monoterpenes showed broader diurnal presence, with early morning and evening peaks in all seasons, and a summer median of 0.51 ± 0.20 ppbv.

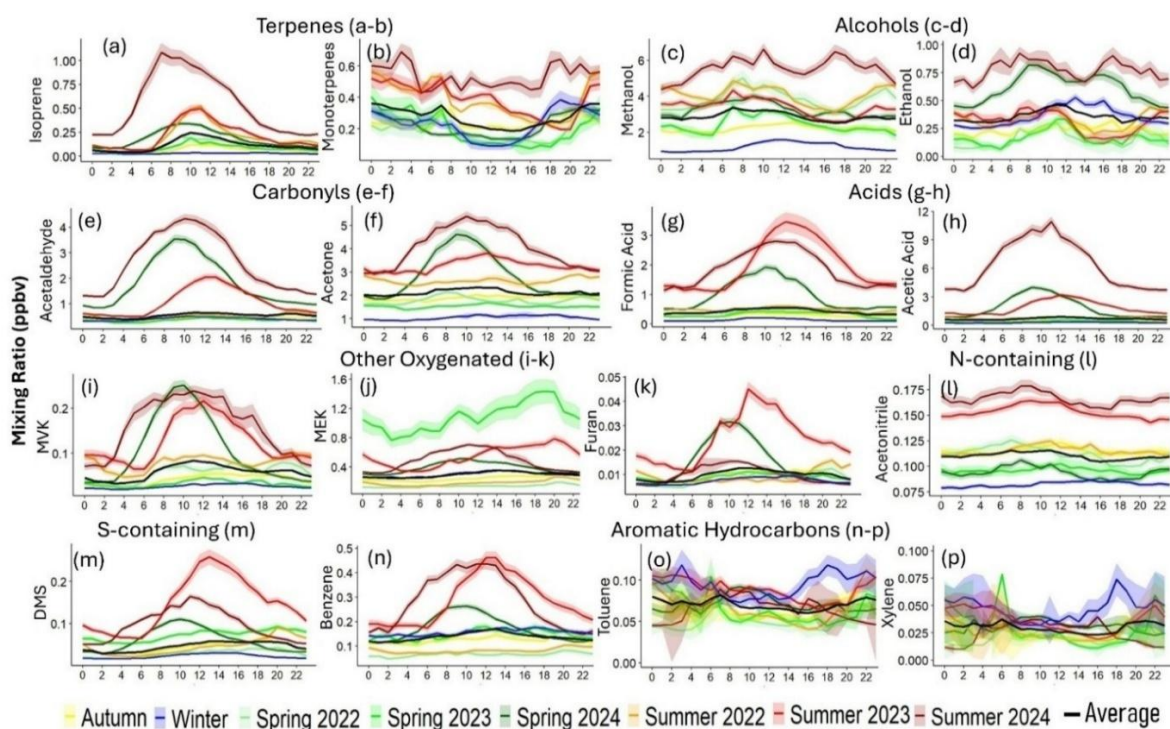


Figure 4: Diurnal profiles (hourly averages) for mixing ratio (ppbv) of major VOCs. It includes terpenes(a-b), oxygenated VOCs (alcohols, carbonyls, acids, other oxygenated (c-k)), nitrogen and sulfur-containing VOCs (l-m), and aromatic hydrocarbons (n-p), measured at a rural background site in Cyprus from 2022 to 2024. Seasonal variations are color-coded, highlighting year-wise spring and summer trends alongside autumn, winter, and overall averages. Shaded areas indicate standard error for median values. X-axis here represents time in hours in UTC. The VOCs selected in this figure serve as representative indicators for their respective chemical families.

Isoprene and monoterpenes exhibited distinctly different temporal patterns, reflecting their contrasting emission drivers and atmospheric dynamics. The seasonal pattern in isoprene align with established findings, where approximately 77% of annual emissions occur in spring (31%) and summer (46%), driven by elevated light intensity and temperature, which activate isoprene synthase enzymes (Fortunati et al., 2008; Monson et al., 1992). Autumn and winter emissions are markedly

425 lower due to reduced photosynthetic activity and enzymatic downregulation (Panopoulou et al., 2020).
 Monoterpenes, while also of biogenic origin mainly, demonstrated a more even seasonal
 distribution, with 38% of levels occurring in summer (0.39 ppbv) and 19-22% across other seasons
 (0.20-0.23 ppbv). This distribution reflects not only temperature dependence but also anthropogenic
 430 and ecological responses to herbivory, wounding, and water stress, particularly relevant under
 changing climate conditions (Bourtsoukidis et al., 2025; Peñuelas and Staudt, 2010; Strada et al.,
 2023; Wang et al., 2022a). The winter levels remain relatively stable, as stored monoterpenes in
 plant tissues continue to be emitted even during periods of low biosynthesis (Bourtsoukidis et al.,
 2024; Ciccioli et al., 2023; Wang et al., 2022a, 2024). Consistent with the findings of Debevec et
 435 al., 2018, our observations also revealed elevated nighttime concentrations of monoterpenes, likely
 attributable to reduced oxidative sinks, boundary layer dynamics, and continuous emissions from
 vegetative storage pools.

3.3.2 Oxygenated VOCs

Methanol and ethanol exhibited moderate diurnal variation with elevated daytime and evening
 440 levels in summer (Fig. 4c-d). Median methanol observed as $\sim 5.52 \pm 1.27$ ppbv and ethanol as
 $\sim 0.76 \pm 0.26$ ppbv in summer 2024, reflecting both biogenic emissions and anthropogenic activities
 such as solvent use and burning. Carbonyls including acetaldehyde (2.38 ± 1.21 ppbv) and acetone
 (3.82 ± 1.02 ppbv), displayed sharp midday peaks in summer, consistent with secondary formation
 via photochemical oxidation. Their diurnal and seasonal consistency between 2023 and 2024
 445 highlights photochemical production pathways. Organic acids such as formic and acetic acid
 showed significant daytime enhancements in summer, particularly in 2024. Their median mixing
 ratios reached 1.76 ± 0.70 ppbv and 5.60 ± 2.75 ppbv, respectively, suggesting combined direct
 emissions and oxidative formation, possibly influenced by biomass burning or regional wildfires.
 Among other OVOCs, methyl vinyl ketone (MVK), a product of isoprene oxidation, peaked at
 450 0.15 ± 0.08 ppbv during summer midday. Furan, a tracer of biomass burning, showed morning
 maxima in summer 2023, indicating episodic fire activity. Consistent with isoprene emission, MVK
 also shows almost 42% of its emission during the summer season. A total of 60-70% levels of acetic
 acid, acetaldehyde, acetone, formic acid, and methanol emissions occur during summer and spring
 season, collectively showing their biogenic origin and secondary formation.

455 3.3.3 Nitrogen and sulfur-containing VOCs

Acetonitrile (C_2H_3N), a N- containing VOC and a biomass burning tracer, showed minimal diurnal
 variation (Fig. 4l) due to its long atmospheric lifetime, which is approximately 620 days (Robles,
 2005). Nevertheless, elevated levels were observed throughout the summer of 2023–2024, pointing
 to regional fire events during summer or long-range transport. The median acetonitrile seasonal
 460 mixing ratios were found to be highest during summer 2024 as 0.17 ± 0.02 ppbv, while lowest during
 winter as 0.08 ± 0.01 ppbv. DMS (C_2H_6S), a S- containing VOC, exhibited its highest average levels
 during summer 2023 (0.12 ± 0.09 ppbv) (Fig. 4m), likely due to enhanced marine biogenic emission

from the nearby Mediterranean sea. The pronounced seasonal variation of DMS supports a temperature-driven biogenic source (Deschaseaux et al., 2022; Edtbauer et al., 2020).

3.3.4 Aromatic hydrocarbons

Benzene showed summer midday enhancements (0.26 ± 0.12 ppbv), while toluene and xylenes exhibited flatter or bimodal patterns (high during morning and evening, probably due to increased vehicular activities) with higher variability. Toluene and xylenes mixing ratios peaked in winter and spring, consistent with local combustion under shallow boundary layers. The large standard error variation (Fig. 4o-p), especially for xylenes and toluene, indicates episodic anthropogenic inputs from traffic or regional pollution. Compared with benzene, these more reactive aromatics exhibited flat or noisy diurnal profiles with limited seasonal variability, aside from the winter-spring xylenes enhancement.

3.4. Temperature-driven enhancement of VOCs emissions in EMME region

3.4.1. Temperature sensitivity of VOCs

The variation in VOCs mixing ratios with temperature revealed clear temperature-dependent trends across most VOCs (Fig. 5). Among terpenes, isoprene, showed strong responses to rising temperatures, as their mixing ratio increased markedly beyond 25 °C, with a peak observed between 35–38 °C (0.53 ± 0.22 ppbv). There is an exponential increase in isoprene levels up to 38 °C, however, a slight decline (0.02–0.06 ppbv) in its level above 38 °C suggests crossing a tipping point through inhibition under extreme heat stress conditions (Fortunati et al., 2008). Monoterpenes exhibited a comparatively low increase in mixing ratios with an increase in temperature, and its trend was more gradual. The moderate rise at 35–38 °C (from 0.34 to 0.41 ppbv, ~21% increase) supports a biogenic contribution during summer, while the relatively stable levels across temperature bins may indicate sustained emissions throughout the day.

DMS levels also rose steadily with temperature, with maximum values observed between 35–38 °C (0.18 ± 0.13 ppbv). This trend is likely driven by enhanced microbial activity in the Mediterranean Sea (Edtbauer et al., 2020) and increased volatilization rates in warm, semi-arid, and coastal conditions typical of the region (Edtbauer et al., 2020).

OVOCs also exhibited strong temperature dependence. Acetone and methanol levels increased progressively, reflecting a combination of direct biogenic emissions and secondary formation via oxidation of precursor compounds (Huang et al., 2020; Liu et al., 2009; Mellouki et al., 2015; Wang et al., 2022b). Methanol levels rose sharply above 38 °C (~19% with mean value of 4.85 ± 0.91 ppbv). Elevated methanol levels at high temperature are primarily consistent with biogenic sources, such as plant metabolism and microbial activity, although emissions from forest fires may also contribute under high-temperature conditions. (Paton-Walsh et al., 2008; Yokelson et al., 1999; Dorokhov et al., 2018). Ethanol also shows temperature dependence, but it is weaker compared to methanol likely reflects its mixed source profile at the

study site. While methanol is largely controlled by temperature-driven biogenic emissions and photochemistry, ethanol receives significant contributions from non-biogenic sources (e.g., solvent use, vehicular exhaust, long-range transport, and agriculture), which are less temperature-sensitive. As shown in Fig. 5, ethanol levels increase markedly only above 38 °C, suggesting additional evaporative emissions from anthropogenic sources. Acetic acid and acetaldehyde showed particularly sharp increases above 35 °C, with acetic acid levels ~2.8 times higher beyond 38 °C (5.4±2.9 ppbv) when compared with the levels at 35-38 °C. These patterns indicate enhanced photochemical production and possible evaporation-driven emissions (Debevec et al., 2017; Xu et al., 2023; Yuan et al., 2024). The increase of OVOCs under warmer conditions (>38 °C) highlights their potentially significant role in SOA formation in the EMME region (Azmi and Sharma, 2023; Pennington et al., 2021).

Among aromatic hydrocarbons, benzene displayed a notable rise above 35 °C (0.35 ppbv), suggesting contributions from evaporate emissions, or long-range transport (Dimitriou and Kassomenos, 2020; Gjesteland et al., 2019). Another possible reason for the high daytime benzene levels, particularly during hot days (AT>35 °C) could be heat-stress-induced biogenic emissions (Misztal et al., 2015), where vegetation was found to release diverse benzenoid compounds under stress, potentially for communication or protective functions. However, the exact cause remains uncertain and warrants further investigation. In contrast, both toluene and xylenes levels were highest at lower temperatures (<15 °C), likely reflecting solvent use, decreased photochemistry, or indoor emissions during cooler periods (Abbasi et al., 2020; Song et al., 2011). A minor elevation (~0.01 ppbv) in toluene and xylenes was observed at 35–38 °C, potentially due to evaporative processes or reduced atmospheric dispersion.

3.4.2. Intensified VOC loading during the Summer 2024 heatwaves

June 2024 was the hottest month on record in Cyprus and coincided with the highest TVOCs mixing ratios observed during the 2022–2024 period (25.76 ± 8.29 ppbv), exceeding summer maxima recorded in 2023 and 2022 (Fig. S4). As illustrated in Fig. 4, most VOCs also showed markedly higher emissions in 2024 compared to previous years. Considering that our observations span only until 15 June 2024, we focused on the comparison of the first half of June across the three years. The data revealed that TVOCs mixing ratio in early June 2024 (23.53 ppbv) were more than double compared to those measured during the same period in 2022 (9.98 ppbv) and 2023 (8.59 ppbv). This sharp rise was accompanied by extreme meteorological conditions: record-breaking average temperatures (24.95 °C), intense SR (314.81 Wm^{-2}), and very low RH (27%). In contrast, temperatures in early June 2022 and 2023 averaged 21.91 °C and 20.61 °C, with RH at 59.73% and 56.39%, respectively.

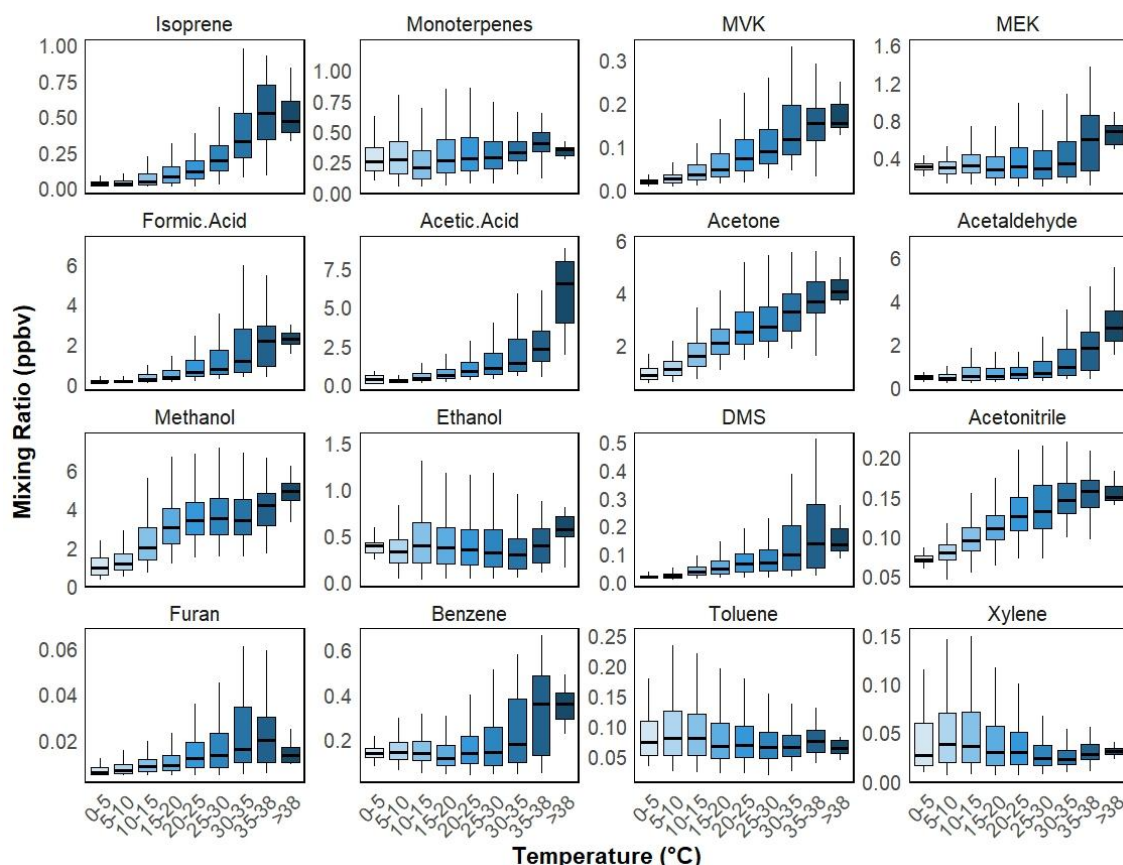


Figure 5: Temperature Dependence of Ambient VOC mixing ratio in Cyprus. Box plots showing the variation in ambient mixing ratio (in ppbv) of selected VOCs across different temperature ranges (°C) at a rural background site in Cyprus. Each box represents the interquartile range (IQR) with the median marked, and whiskers indicating variability outside the upper and lower quartiles. The VOCs selected in this figure are serve as representative indicators for their respective chemical families.

These conditions favored both primary biogenic emissions and secondary atmospheric processes. Rapid vegetative growth in spring, followed by early thermal stress, likely stimulated isoprene and monoterpene emissions, and maybe also benzene. Elevated temperatures and oxidant availability enhanced both the emission (Bourtsoukidis et al., 2014) and formation of OVOCs such as acetaldehyde, acetic acid, and methanol. Concurrently, increased levels of CO (144.77 ppbv) and O₃ (58.77 ppbv) in 2024, compared to 2022 (132.69 and 45.13 ppbv) and 2023 (119.27 and 51.88 ppbv), indicate stronger photochemical activity and increased ozone formation potential (OFP).

Midday peaks in isoprene, along with enhanced OVOCs mixing ratios (acetone, acetic acid, acetaldehyde), reflect intense photochemical production under elevated SR. Also, the elevated nighttime MVK/Isoprene ratio (0.96) indicates substantial isoprene oxidation, consistent with an aged air mass that has undergone prolonged exposure to OH radicals. This further supports enhanced oxidation of biogenic precursors (Guo et al., 2012). In contrast, the lower daytime ratio (0.62) suggests comparatively fresher isoprene emissions, reflecting active daytime production of isoprene and limited conversion to MVK, which requires time to form through photochemical processes (Apel et al., 2002). The findings suggest that escalating heatwave intensity and frequency in the EMME region could significantly increase VOC burdens, contributing to secondary pollutant formation (O₃ and SOA) and worsening regional air quality.

Wildfire activity may also exhibit pronounced variability and likely influenced the observed VOC composition at the CAO-AMX site (Fig. S6). From 2022-2024, the regional fire activity began in spring; however, its timing and intensity differed substantially. In 2022, the burned area increased rapidly in early summer, exceeding ~1800 ha in June (EEFIS, 2025). In 2023, fire activity was comparatively weaker during spring and early summer, with the largest burned areas occurring later in the season (August-September, 900 and 700 ha respectively). In contrast, 2024 was characterized by an earlier onset of wildfires, with fires occurring from April onwards and peak burned areas of >700 ha in May and ~2250 ha in June. As a result, springtime fire influence was strongest in 2024 compared to 2022 and 2023 (EEFIS, 2025). These spring and summer wildfire episodes, particularly in 2024, are expected to have enhanced biomass-burning-related VOCs at the site. Fire-influenced air masses typically exhibit elevated concentrations of acetonitrile, methanol, and other OVOCs.

3.5. Inter-species relationship

Based on the comprehensive correlation analysis in Fig. 6, clear patterns emerge among VOCs, meteorological parameters, and pollutant levels. The overall correlation matrix reveals strong positive associations among OVOCs, including acetaldehyde, acetic acid, acetone, methanol, and formic acid ($r > 0.6-0.9$), suggesting common sources and secondary formation. Ethanol does not cluster closely ($r < 0.5$) with other OVOCs in the hierarchical analysis (Fig. S7). Its more mixed source profile, including contributions from both local and transported anthropogenic activities, results in distinct temporal behaviour relative to primarily biogenic or photochemical VOCs. Thus, the clustering and weaker temperature dependence are consistent with ethanol's more complex source influences at the site. Isoprene strongly correlates with MVK ($r = 0.75$), indicating secondary photochemical formation and biogenic influence during high radiation periods. Isoprene is also positively associated with temperature, and solar radiation ($r = 0.5$) indicating that variability in isoprene and MVK is driven by photochemical processing. On the contrary, monoterpenes show weak correlation at both interspecies levels, and with meteorological parameters, showing its mixed sources from both biogenic and anthropogenic origins.

Aromatic hydrocarbons (e.g., toluene and xylenes) form another cluster with moderate-to-strong correlations ($r = 0.94$), consistent with urban combustion sources. Among aromatic hydrocarbons, benzene show comparatively weaker relation with toluene and xylenes. Benzene-toluene ($r = 0.6$) and benzene-xylenes ($r = 0.4$) correlation is moderate in winter, maybe due to low photochemical loss due to low temperature, but weakens in summer ($r < 0.3$) due to faster toluene degradation and variable sources (Gelencsér et al., 1997). Toluene is showing weak correlation ($r < 0.3$) with other VOCs and environmental parameters, showing their main source is anthropogenic in origin. Whereas benzene shows strong correlation with other VOCs specially OVOCs ($r > 0.6$) showing either similar sources or evaporation process.

At the rural Cyprus background site, CO shows a diurnal cycle with an early-morning minimum and an afternoon-late evening maximum, which may be driven by boundary-layer dilution after sunrise and subsequent accumulation under regionally influenced air masses. The generally weak correlation between CO and aromatic VOCs suggests that CO variability is primarily controlled by

regional transport and its longer atmospheric lifetime, rather than local traffic emissions. Seasonally, CO exhibits moderate correlations with benzene in autumn ($r \approx 0.4$) and winter ($r \approx 0.5$), likely reflecting enhanced regional combustion sources, reduced boundary-layer heights, and slower photochemical removal during colder months. Correlations are weak in spring and summer due to stronger photochemical processing and deeper boundary layers. CO shows no correlation with toluene or xylene in any season, consistent with their shorter lifetimes and the absence of fresh local emissions at this background site.

RH shows negative correlations with most VOCs, likely due to emission suppression or enhanced wet removal. O₃ correlated moderate but positive ($r > 0.25$) with methanol, acids, acetone, isoprene and also with AT and SR reflecting active photochemical processes especially during summer. Although in correlation matrix, WS and WD show weaker correlations but may still influence variability, as suggested by backward trajectory analysis in section 3.6. Overall hierarchical clustering in Fig. 6 groups of species by chemical and functional similarity. OVOCs cluster tightly, as do aromatics, while temperature, radiation, and ozone form a separate group. These clusters reflect common sources and atmospheric behaviors. Seasonal correlation matrices (Fig. S7) also highlight dynamic VOC-environment interactions, with the strongest correlation among most VOCs during summer.

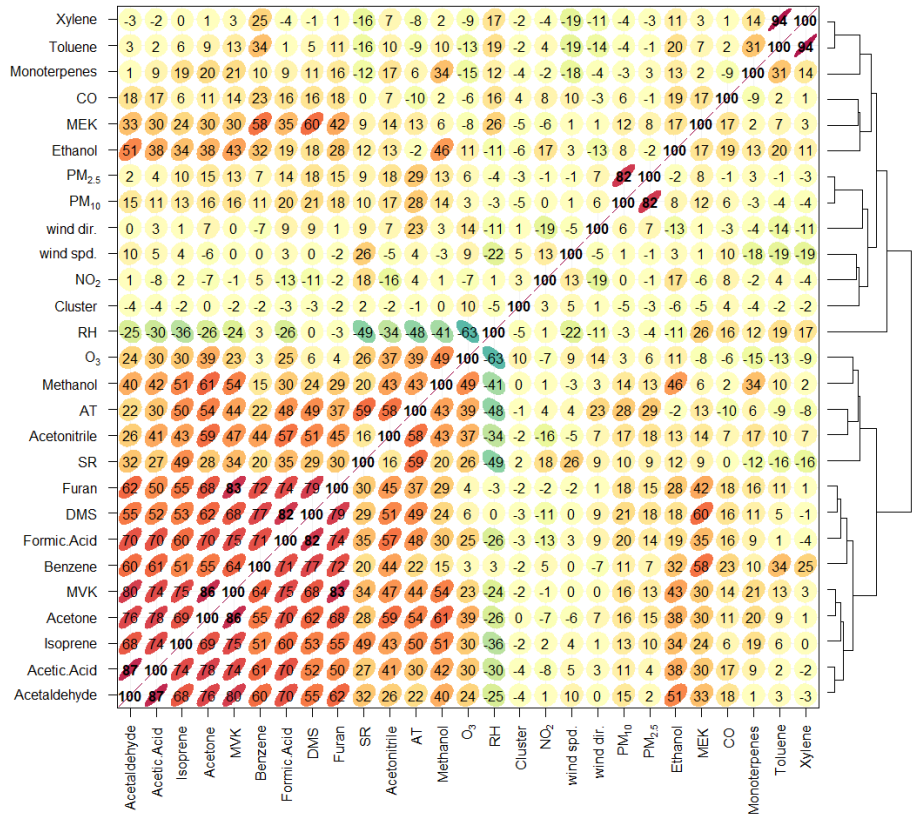


Figure 6: Correlation matrix and hierarchical clustering of VOCs, meteorological parameters, and air pollutants. Pearson correlation matrix illustrates the relationships among VOCs, meteorological variables (e.g., temperature, solar radiation, humidity), and air pollutants (NO₂, O₃, CO, PM_{2.5}, PM₁₀). Here, cluster refers to the air-mass trajectory cluster index obtained from the HYSPLIT back-trajectory analysis (shown in Fig. 7), where each number corresponds to a distinct air-mass transport pathway. Positive correlations are shown in orange to red, while negative correlations are represented in green to blue. Hierarchical clustering (dendrogram) on the right groups variables with similar correlation patterns.

The diurnal variation of selected interspecies mixing ratios (Fig. S8) reveals distinct patterns driven by emission sources, photochemical processing, and atmospheric reactivity. Ratios such as toluene/benzene, and xylenes/benzene decrease during the day (minima: ~0.45, and ~0.15 respectively), indicating stronger photochemical degradation of toluene and xylenes compared to the more stable benzene (Tiwari et al., 2010). The MVK/isoprene ratio increases from ~0.5 in the early morning to ~1.0 by midday, reflecting active daytime oxidation of isoprene to MVK (Guo et al., 2012). Similarly, the isoprene/monoterpenes ratio peaks around 1.2 between 10:00–14:00 UTC, highlighting the temperature- and radiation-driven nature of isoprene emissions, while monoterpenes remain relatively stable (Kiendler-Scharr et al., 2009). The ozone/NO₂ ratio exhibits a strong daytime increase, rising from ~30 to over 60 at noon, reflecting photochemical ozone formation from NO₂ under solar radiation. Overall, the interspecies ratios effectively differentiate between primary emissions and secondary formation, with photochemically active species (isoprene, MVK, ozone) showing strong diurnal variability.

3.6. Air mass origins and transport patterns

Using the HYSPLIT model to calculate air mass backward trajectories, we identified seven air mass clusters (Fig. 7) influencing VOCs transport and composition over Cyprus. The dominant clusters originated from Northwest Asia (C4, 34.3%) and Europe (C3, 33.6%), jointly accounting for more than two-thirds of the long-range air mass transport. These clusters represent strong, recurrent long-range synoptic-scale transport from urbanized and industrialized continental regions, which occur more frequently than locally influenced trajectories, and thus play a key role in shaping seasonal and spatial VOC variability. Marine-influenced air masses (C2, 13.4%) reflected mixed characteristics, carrying both marine biogenic signatures and aged anthropogenic pollution. Less frequent clusters included air masses from North Africa (C1, 8.1%), Western Turkey (C5, 4.3%), Middle East (C6, 4.3%), and stagnant local conditions (C0, 1.7%). Notably, C6, although infrequent overall, was more prevalent during winter (10.7%) and exhibited elevated levels of benzene, toluene, and xylenes (BTX) during winter, suggesting long-range transport of aromatic hydrocarbons from the Middle East. C0, characterized by low wind speeds and residence near the site, showed high concentrations of primary pollutants such as benzene, acetonitrile, ethanol, and NO₂, consistent with local anthropogenic and biomass burning sources. Despite dominant local Northwest winds (as shown in Windrose diagram Fig. 2d), back trajectories show multi-day synoptic transport. The more frequent large-scale flows (C3-C4) therefore contribute over two-thirds of long-range air mass transport, exceeding the influence of local pathways (C1-C2). Distinct chemical signatures were observed across clusters. European clusters (C3) exhibited elevated OVOCs such as methanol, acetone, and acetic acid, indicating the presence of aged air masses and secondary formation processes. C3 also carried higher levels of isoprene and MVK, possibly from transported biogenic emissions. C4 was associated with elevated O₃, potentially due to photochemical aging during long-range transport. In contrast, Western Turkey (C5) air masses had generally low VOC mixing ratios, though slight O₃ enhancement was observed, possibly due to regional NO_x-rich emissions and transport.

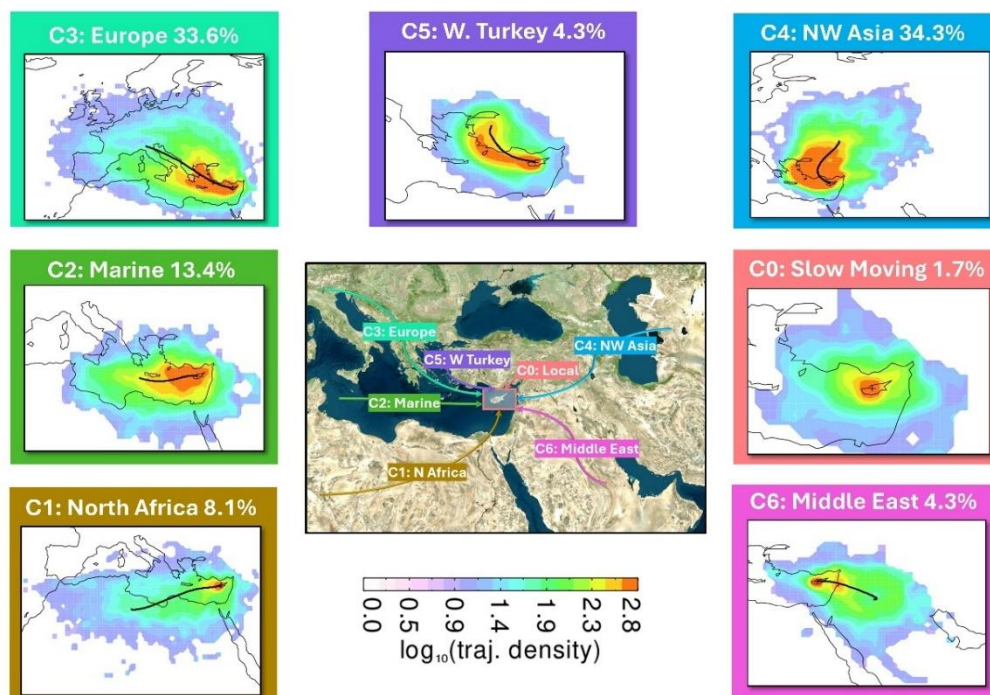


Figure 7: Five-day air mass backward trajectories starting at 500 m above the ground level starting at every hour. These were calculated from the NOAA ARL PC-version HYSPLIT transport and dispersion model using gridded wind fields from the GDAS, which has a spatial resolution of $1^\circ \times 1^\circ$ longitude by latitude and a time resolution of 1 hr (Draxler and Hess, 1999) to demonstrate the influence of long-range transported air masses at the measurement site.

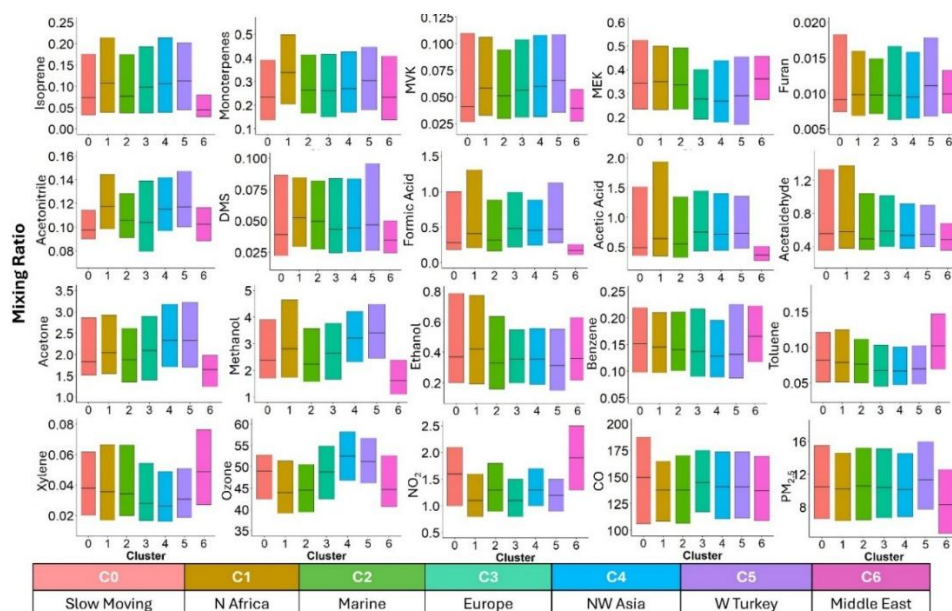


Figure 8: Mixing ratios of VOCs, and other air pollutants (in ppbv except $\text{PM}_{2.5}$, which is in $\mu\text{g m}^{-3}$) across air mass clusters in the regional site at Eastern Mediterranean. Boxplots showing the distribution of mixing ratios of selected VOCs, and other air pollutants for different air mass clusters (C0–C6) identified by back trajectory analysis. The black line within each box represents the median; boxes indicate the interquartile range, and whiskers denote variability outside the upper (75th) and lower (25th) quartiles. The VOCs selected in this figure are serve as representative indicators for their respective chemical families.

Seasonal analysis (Fig. S9-S10) revealed pronounced variability in both air mass distribution and VOC loading. Spring and summer were dominated by clusters from Europe and NW Asia, corresponding with enhanced biogenic emissions, active photochemistry, and increased long-range transport. These seasons showed elevated mixing ratios of isoprene, monoterpenes, methanol, acetic acid, and O₃. Summer and spring exhibited the highest overall VOC and O₃ levels across most clusters, underscoring strong atmospheric photochemical reactivity. Summer maintained high VOC loadings, particularly in C2 and C3 clusters, driven by elevated temperatures and insolation. Autumn displayed a relatively balanced air mass distribution with moderate VOC levels. Winter was characterized by reduced photochemical activity and lower overall VOC concentrations, except for C6, which exhibited persistently elevated anthropogenic markers such as BTX likely associated with heating-related emissions and regional transport under more stagnant meteorological conditions. These findings underscore the interplay between long-range transport, regional meteorology, and local emissions in shaping atmospheric VOCs variability over Cyprus. The combined influence of continental, marine, and regional sources contributes to complex spatiotemporal variability, with implications for air quality, ozone formation, and atmospheric oxidation capacity in the EMME region.

3.7. WRF-Chem model simulations

WRF-Chem simulations were evaluated against high-resolution PTR-ToF-MS observations at CAO-AMX across multiple seasons to assess the model's capability to reproduce VOCs variability in the EMME region. BVOCs (isoprene and monoterpenes) emissions were online estimated by MEGAN v2.1 (Guenther et al., 2012), while anthropogenic emissions were derived from EDGAR-HTAP for the outer domains and a Cyprus high-resolution inventory for the innermost domain (Georgiou et al., 2020, 2022). Key biogenic species include isoprene and monoterpenes, while anthropogenic aromatics include benzene, toluene, and xylenes. These species, together with OVOCs, are evaluated using seasonal distributions, diurnal behavior, and annual averages (Fig. 9, Figs. S11–S12). The model successfully captured the seasonal and diurnal variability of several VOCs, particularly the strong summer enhancement associated with elevated temperatures and photochemical activity. Terpenes exhibited pronounced daytime and summertime maximum levels in both observations and simulations, consistent with temperature- and light-driven biogenic emissions reported at other Mediterranean and European background sites (Ge et al., 2024; Genard-Zielinski et al., 2015). Among the evaluated species, **isoprene showed the best agreement**, with the model reproducing its temporal variability ($r = 0.52$, $p < 0.05$), although absolute mixing ratios were overestimated by approximately a factor of three. This overestimation is consistent with previous European-scale studies showing that MEGAN-based WRF-Chem simulations tend to overpredict isoprene emissions, particularly in southern and semi-arid regions (Morichetti et al., 2022). These biases have been primarily attributed to the model's lack of an effective drought stress and soil-moisture constraints in the current MEGAN emission parameterization (Ge et al., 2024; Genard-Zielinski et al., 2015; Morichetti et al., 2022).

715 In contrast, Strada et al. (2023) reported strong underprediction of isoprene mixing ratios for Cyprus, that was also attributed to uncertainties in water availability dynamics, demonstrating inadequate understanding of the role of soil moisture in regulating isoprene emissions in the EMME region. These contrasting results emphasize the need for improved, region-specific parameterizations of plant stress responses under extreme meteorological conditions. For monoterpenes, the model

720 significantly underestimated the measured mixing ratios, by up to a factor of 6-7 during daytime and 3-4 at night (Fig. S12). The underestimation of monoterpenes in the model could stem from the assumption that their emissions are primarily controlled by light and temperature. If monoterpenes originate predominantly from storage pools, their fluxes will exhibit only a weak light dependency, which could explain the observed discrepancies. In contrast, the overestimation of isoprene in our

725 study may be linked to heat- and drought-induced stress during the summer months. Such conditions can alter plant carbon allocation strategies, leading to reduced isoprene emissions and enhanced monoterpene release, consistent with observations reported by Byron et al. (2022), but this mechanism may not be represented in current MEGAN parameterizations. Incorporating storage-driven emissions and stress-dependent emission pathways would therefore be critical for improving monoterpene simulations in hot, semi-arid regions. Biogenic emission (isoprene, and monoterpenes) parameterisations require improved representation of high-temperature and water-stress responses (Sindelarova et al., 2014). In particular, heat-stress emission factors and land-atmosphere coupling should be targeted in future work, consistent with studies showing that drought stress algorithms improve the capability of MEGAN isoprene emission estimates (Wang et al., 2022c).

730 Photochemical reaction pathways, chemical ageing and deposition processes that show sensitivity to heat wave atmospheric states can be investigated to more properly quantify atmospheric removal in WRF-CHEM (Knote et al., 2015a,b). In addition, wildfire and anthropogenic VOC emissions should be produced using high spatiotemporal (hourly, 1x1 km) resolution to capture the impact of applying diurnal variability and vertical distribution to fire emissions, along with VOC speciation

735 that better reflects regional source profiles.

740 OVOCs including acids, aldehydes, and ketones showed strong diurnal and seasonal enhancements during summer, consistent with intense photochemical production and evaporative processes, yet were underestimated by factors of 3-12 depending on chemical class. These discrepancies are aligned with underprediction of OVOCs across Europe due to missing secondary formation chemistry, incomplete representation of oxidation products, and inadequate representation of

745 temperature-dependent evaporative emissions (Ge et al., 2024). Taken together, these results indicate that there are significant deficiencies in the representation of OVOCs precursors and multi-phase processing by RACM.

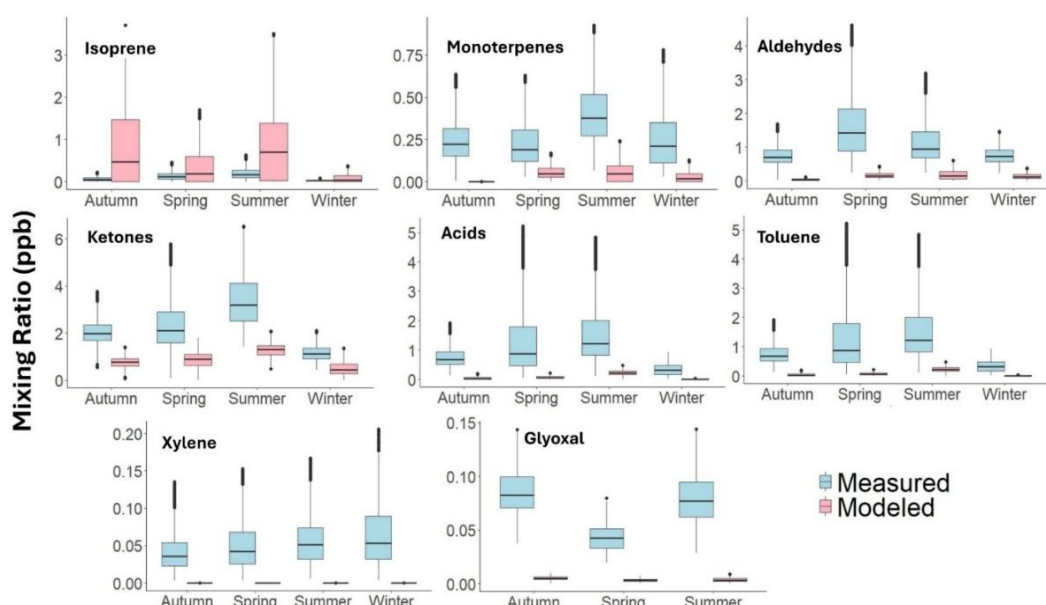


Figure 9: Seasonal variation among measured and simulated VOCs mixing ratios at rural background site in Cyprus. The selection, grouping, and classification of VOCs presented in this figure follow the compound categories available within the chemical mechanism of the model to ensure consistency between observations and simulations.

Aromatic hydrocarbons like toluene and xylenes showed relatively flat diurnal profiles with slight morning and evening peaks in cooler months (Fig. S12), indicative of local anthropogenic sources under stable atmospheric conditions. However, WRF-Chem significantly underestimated these species, particularly xylenes, implying incomplete representation of urban emissions, traffic sources, and solvent use in anthropogenic inventories. Glyoxal, an important secondary oxidation product, exhibited a clear midday peak in measurements, but modeled values were significantly lower, even though the timing of the peak was captured, suggesting underestimation of precursor VOCs and low yields in photochemical pathways.

Seasonal comparisons (Fig. 9) confirmed these discrepancies. Observed levels of isoprene, aldehydes, ketones, and monoterpenes peaked in spring and summer, consistent with enhanced biogenic and photochemical activity. Modeled outputs, however, underrepresented OVOCs and AVOCs across all seasons. Despite some agreement in seasonal patterns, WRF-Chem was unable to reproduce the magnitude and timing of peak VOC levels. The annual comparison (Fig. S11) showed that while isoprene mean simulated values were relatively aligned with observations, most other VOC groups, particularly OVOCs and aromatics, were systematically underestimated.

EDGAR generally reports higher VOC emissions compared to EMEP or CAMS-REG-AP due to differences in industrial combustion and fugitive sector representation (Thunis et al., 2021). However, these differences in VOC emissions have a limited impact on European-scale ozone modelling, and chemical regime and meteorology remain dominant drivers of O_3 variability (Thunis et al., 2021). A high-resolution Cyprus inventory may significantly improve the spatial representation of anthropogenic emissions and reduces NO_2 and O_3 biases compared to simulations based on CAMS (Georgiou et al., 2022), but uncertainties in BVOCs and OVOCs sources remain the largest contributors to discrepancies.

Overall, the analysis reveals that the heat extremes of the Eastern Mediterranean evoke a distinctive VOC emission response that enhances differences between observations and models based on traditional parameterizations developed in temperate conditions. The systematic underprediction of monoterpenes and OVOCs, when taken together with the overestimation of isoprene, suggests that incorporation of drought-sensitive plant physiology, storage-driven emission dynamics, and extended secondary oxidation chemistry within regional air quality models is required. Both focused biogenic emission modeling efforts and OVOC reaction pathways are necessary to appropriately quantify atmospheric reactivity and chemical transport in this climate-sensitive region. Also, as the frequency and intensity of heatwaves are projected to increase under future climate scenarios (Zittis et al., 2021; Lelieveld et al., 2016), the non-linear response of VOC emissions to temperature stress must be better parameterized to prevent significant underestimations of SOA and ozone formation.

4 Conclusions

This study provides the first multi-year, high-resolution characterization of volatile organic compounds (VOCs) in the Eastern Mediterranean based on continuous PTR-ToF-MS measurements at the Cyprus Atmospheric Observatory. By integrating detailed chemical observations with meteorological analyses, HYSPLIT air-mass trajectories, and WRF-Chem simulations, we offer a comprehensive assessment of the processes governing VOC variability in a rapidly warming and climatically sensitive region. Across the 2022–2024 period, oxygenated VOCs dominated the total VOC burden (~79%), reflecting the strong influence of direct sources, secondary formation, and long-range transport on regional atmospheric composition.

Among terpenes, isoprene, exhibited pronounced summer peaks and daytime maxima, closely linked to solar radiation and temperature. Isoprene increased with temperature up to 38 °C before declining, suggesting thermal stress emission suppression, while monoterpenes showed both daytime and nocturnal presence may be due to diverse emission drivers from vegetation, boundary layer dynamics, or anthropogenic contribution. OVOCs (acetone, methanol, acetic acid, acetaldehyde) were enhanced during hot, dry periods, indicating amplified primary emissions and secondary formation. DMS peaks under high temperatures pointed to microbial activity and volatilization in semi-arid and coastal environments. Among aromatic hydrocarbons benzene increased sharply above 35 °C, likely from biogenic origin, evaporative emissions and long-range transport, whereas toluene and xylenes were more abundant in cooler periods and degraded rapidly at higher temperatures.

HYSPLIT trajectory cluster analysis revealed elevated aromatic hydrocarbons, under the influence of long-range transported continental air masses from middle east cluster (C6). The variability in organic acid and sulfur compound was attributed to marine and North African air masses. Slow-moving local air masses, although infrequent, showed disproportionately high VOCs mixing ratio, indicating strong local emission influence. Seasonal patterns showed summer dominated by biogenic emissions and active photochemistry, while winter was characterized by anthropogenic influences under stagnant conditions; spring emerged as a particularly reactive period.

815 WRF-Chem model simulations captured general seasonal patterns of isoprene but systematically underpredicted OVOCs, aromatics, and secondary VOCs, with notable temporal mismatches in diurnal cycles. These shortcomings reflect limitations in emission inventories, oxidation schemes, and meteorological coupling, particularly for region-specific and stress-induced emissions.

820 Overall, this work demonstrates that VOC variability in the Eastern Mediterranean is governed by the combined influence of extreme heat, seasonal variation, regional transport, and complex photochemistry. The findings underscore the need for improved VOC parameterizations in regional to global models, region-specific source inventories, and adaptive air quality management strategies that account for both local and long-range transported emissions. With heatwaves expected to intensify across the EMME region, understanding how VOC emissions respond to climatic extremes is critical for predicting future ozone formation, secondary organic aerosol production, and the broader oxidative capacity of the atmosphere. This multi-year dataset provides a valuable foundation for future source apportionment analysis, evaluation of regional model performance, and development of improved emission inventories. As one of the few long-term VOC records in the region, it represents an important step toward constraining atmospheric processes in a globally significant climate hotspot.

Data availability. All data used in this study are openly available at <https://zenodo.org/records/17483541>

Author Contribution.

835 AG: Conceptualization, Data curation, Methodology, Validation, Formal analysis, Investigation, Writing – original draft, Writing – review & editing, Visualization. MD: Data curation, Methodology, Software, Validation, Formal analysis, Writing – review & editing. AC: Methodology, Data curation, Formal analysis, Writing – review & editing. TC: Data curation, Software, Writing – review & editing. VPK: Methodology, Software, Writing – review & editing. CS: Data curation, Resources, Writing – review & editing. MV: Methodology, Writing – review & editing. SN: Formal analysis, Writing – review & editing. TJ: Writing – review & editing. JB: Writing – review & editing. JW: Writing – review & editing. NM: Writing – review & editing. EL: Writing – review & editing. JS: Funding acquisition, Resources, Writing – review & editing. EB: Conceptualization, Project administrator, Methodology, Validation, Investigation, Writing – review & editing, Supervision.

Competing interests.

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

Acknowledgements.

This work was supported by the European Union's Horizon 2020 research and innovation program (grant No. 856612) and the Cyprus Government (EMME-CARE). The authors thank the CAO technical team for the routine upkeep of the AMX station. EB and SN acknowledge support from the project EVOPOLIS funded by The Cyprus Research and Innovation Foundation (RIF), implemented under the Cohesion Policy Programme “Thalia 2021-2027” and co-funded by the EU. TJ and VPK are supported by the European Union ERC-2022-STGERC-BAE-Project: 101076311. Views and opinions expressed are however those of the author(s) only and do not necessarily reflect those of the European Union or the European Research Council Executive Agency. Neither the European Union nor the granting authority can be held responsible for them.

References

- Abbasi, F., Pasalari, H., Delgado-Saborit, J. M., Rafiee, A., Abbasi, A., and Hoseini, M.: Characterization and risk assessment of BTEX in ambient air of a Middle Eastern City, Process Safety and Environmental Protection, 139, 98–105, <https://doi.org/10.1016/j.psep.2020.03.019>, 2020.
- Apel, E. C., Riemer, D. D., Hills, A., Baugh, W., Orlando, J., Faloon, I., et al.: Measurement and interpretation of isoprene fluxes and isoprene, methacrolein, and methyl vinyl ketone mixing ratios at the PROPHET site during the 1998 intensive, J. Geophys. Res.-Atmos., 107, ACH-7, <https://doi.org/10.1029/2000JD000225>, 2002.
- Atkinson, R.: Atmospheric chemistry of VOCs and NO_x, Atmos Environ, 34, 2063–2101, [https://doi.org/10.1016/S1352-2310\(99\)00460-4](https://doi.org/10.1016/S1352-2310(99)00460-4), 2000.
- Azmi, S. and Sharma, M.: Global PM_{2.5} and secondary organic aerosols (SOA) levels with sectorial contribution to anthropogenic and biogenic SOA formation, Chemosphere, 336, <https://doi.org/10.1016/j.chemosphere.2023.139195>, 2023.
- Baalbaki, R., Pikridas, M., Jokinen, T., Laurila, T., Dada, L., Bezantakos, S., Ahonen, L., Neitola, K., Maissner, A., Bimenyimana, E., Christodoulou, A., Unga, F., Savvides, C., Lehtipalo, K., Kangasluoma, J., Biskos, G., Petäjä, T., Kerminen, V.-M., Sciare, J., and Kulmala, M.: Towards understanding the characteristics of new particle formation in the Eastern Mediterranean, Atmos Chem Phys, 21, 9223–9251, <https://doi.org/10.5194/acp-21-9223-2021>, 2021.
- Bourtsoukidis, E., Williams, J., Kesselmeier, J., Jacobi, S., and Bonn, B.: From emissions to ambient mixing ratios: online seasonal field measurements of volatile organic compounds over a Norway spruce-dominated forest in central Germany, Atmos Chem Phys, 14, 6495–6510, <https://doi.org/10.5194/acp-14-6495-2014>, 2014.
- Bourtsoukidis, E., Behrendt, T., Yañez-Serrano, A. M., Hellén, H., Diamantopoulos, E., Catão, E., Ashworth, K., Pozzer, A., Quesada, C. A., Martins, D. L., Sá, M., Araujo, A., Brito, J., Artaxo, P., Kesselmeier, J., Lelieveld, J., and Williams, J.: Strong sesquiterpene emissions from Amazonian soils, Nat Commun, 9, 2226, <https://doi.org/10.1038/s41467-018-04658-y>, 2018.
- Bourtsoukidis, E., Ernle, L., Crowley, J. N., Lelieveld, J., Paris, J.-D., Pozzer, A., Walter, D., and Williams, J.: Non-methane hydrocarbon (C₂–C₈) sources and sinks around the Arabian Peninsula, Atmos Chem Phys, 19, 7209–7232, <https://doi.org/10.5194/acp-19-7209-2019>, 2019.
- Bourtsoukidis, E., Pozzer, A., Williams, J., Makowski, D., Peñuelas, J., Matthaios, V. N., Lazoglou, G., Yañez-Serrano, A. M., Lelieveld, J., Ciais, P., Vrekoussis, M., Daskalakis, N., and Sciare, J.: High temperature sensitivity of monoterpene emissions from global vegetation, Commun Earth Environ, 5, 23, <https://doi.org/10.1038/s43247-023-01175-9>, 2024.
- Bourtsoukidis, E., Guenther, A., Wang, H., Economou, T., Lazoglou, G., Christodoulou, A., Christoudias, T., Nölscher, A., Yañez-Serrano, A. M., and Peñuelas, J.: Environmental Change Is Reshaping the Temperature Sensitivity of Sesquiterpene Emissions and Their Atmospheric Impacts, Glob Chang Biol, 31, <https://doi.org/10.1111/gcb.70258>, 2025.
- Brito, J., Freney, E., Dominutti, P., Borbon, A., Haslett, S. L., Batenburg, A. M., Colomb, A., Dupuy, R., Denjean, C., Burnet, F., Bourriane, T., Deroubaix, A., Sellegri, K., Borrmann, S., Coe, H., Flamant, C., Knippertz, P., and Schwarzenboeck, A.: Assessing the role of anthropogenic and biogenic sources on PM_{2.5} over southern West Africa

using aircraft measurements, *Atmos Chem Phys*, 18, 757–772, <https://doi.org/10.5194/acp-18-757-2018>, 2018.

900 Bryant, D. J., Nelson, B. S., Swift, S. J., Budisulistiorini, S. H., Drysdale, W. S., Vaughan, A. R., Xu, L., Rickard, A. R., Hewitt, C. N., Nemitz, E., Mullinger, N., Acton, W. J. F., Young, D. E., Allan, J., & Hamilton, J. F.: Biogenic and anthropogenic sources of isoprene and monoterpenes and their secondary organic aerosol in Delhi, India, *Atmos. Chem. Phys. Discuss.*, 2022, 1–35, <https://doi.org/10.5194/acp-2022-xxx>, 2022.

905 Byron, J., Kreuzwieser, J., Purser, G., van Haren, J., Ladd, S. N., Meredith, L. K., Werner, C., and Williams, J.: Chiral monoterpenes reveal forest emission mechanisms and drought responses, *Nature*, 609, 307–312, <https://doi.org/10.1038/s41586-022-05020-5>, 2022.

910 Capes, G., Murphy, J. G., Reeves, C. E., McQuaid, J. B., Hamilton, J. F., Hopkins, J. R., Crosier, J., Williams, P. I., and Coe, H.: Secondary organic aerosol from biogenic VOCs over West Africa during AMMA, *Atmos. Chem. Phys.*, 3841–3850 pp., 2009.

Ciccioli, P., Silibello, C., Finardi, S., Pepe, N., Ciccioli, P., Rapparini, F., Neri, L., Fares, S., Brilli, F., Mircea, M., Magliulo, E., and Baraldi, R.: The potential impact of biogenic volatile organic compounds (BVOCs) from terrestrial vegetation on a Mediterranean area using two different emission models, *Agric For Meteorol.*, 328, 109255, <https://doi.org/10.1016/j.agrformet.2022.109255>, 2023.

915 Civan, M. Y., Elbir, T., Seyfioglu, R., Kuntasal, Ö. O., Bayram, A., Doğan, G., Yurdakul, S., Andiç, Ö., Müezzinoğlu, A., Sofuoglu, S. C., Pekey, H., Pekey, B., Bozlaker, A., Odabasi, M., and Tuncel, G.: Spatial and temporal variations in atmospheric VOCs, NO₂, SO₂, and O₃ concentrations at a heavily industrialized region in Western Turkey, and assessment of the carcinogenic risk levels of benzene, *Atmos Environ*, 103, 102–113, <https://doi.org/10.1016/j.atmosenv.2014.12.031>, 2015.

920 Cyprus Mail: Last month hottest June ever in Cyprus., *Cyprus Mail*, 2024.

925 Das, A., Giri, B. S., and Manjunatha, R.: Systematic review on benzene, toluene, ethylbenzene, and xylene (BTEX) emissions; health impact assessment; and detection techniques in oil and natural gas operations, *Environmental Science and Pollution Research*, 32, 1–22, <https://doi.org/10.1007/s11356-024-35698-1>, 2024.

930 Debevec, C., Sauvage, S., Gros, V., Sciare, J., Pikridas, M., Stavroulas, I., Salameh, T., Leonardis, T., Gaudion, V., Depelchin, L., Fronval, I., Sarda-Esteve, R., Baisnée, D., Bonsang, B., Savvides, C., Vrekoussis, M., and Locoge, N.: Origin and variability in volatile organic compounds observed at an Eastern Mediterranean background site (Cyprus), *Atmos Chem Phys*, 17, 11355–11388, <https://doi.org/10.5194/acp-17-11355-2017>, 2017.

935 Debevec, C., Sauvage, S., Gros, V., Sellegri, K., Sciare, J., Pikridas, M., Stavroulas, I., Leonardis, T., Gaudion, V., Depelchin, L., Fronval, I., Sarda-Esteve, R., Baisnée, D., Bonsang, B., Savvides, C., Vrekoussis, M., and Locoge, N.: Driving parameters of biogenic volatile organic compounds and consequences on new particle formation observed at an eastern Mediterranean background site, *Atmos Chem Phys*, 18, 14297–14325, <https://doi.org/10.5194/acp-18-14297-2018>, 2018.

940 Department of Meteorology (Cyprus). Ministry of Agriculture, Rural Development and Environment, (2024, 15 July). *Monthly Weather Report: The Weather in June 2024* [PDF]. https://www.dom.org.cy/CLIMATOLOGY/English/Monthly%20Weather%20Bulletin/2024/06_Monthly%20Weather%20Report_Jun2024_ENG.pdf

945 Derstroff, B., Hüser, I., Bourtsoukidis, E., Crowley, J. N., Fischer, H., Gromov, S., Harder, H., Janssen, R. H. H., Kesselmeier, J., Lelieveld, J., Mallik, C., Martinez, M., Novelli, A., Parchatka, U., Phillips, G. J., Sander, R., Sauvage, C., Schuladen, J., Stönnner, C., Tomsche, L., and Williams, J.: Volatile organic compounds (VOCs) in photochemically aged air from the eastern and western Mediterranean, *Atmos Chem Phys*, 17, 9547–9566, <https://doi.org/10.5194/acp-17-9547-2017>, 2017.

950 Deschaseaux, E. S. M., Swan, H. B., Maher, D. T., Jones, G. B., Schulz, K. G., Koveke, E. P., Toda, K., and Eyre, B. D.: The Interplay Between Dimethyl Sulfide (DMS) and Methane (CH₄) in a Coral Reef Ecosystem, *Front Mar Sci*, 9, <https://doi.org/10.3389/fmars.2022.910441>, 2022.

Desservettaz, M., Pikridas, M., Stavroulas, I., Bougiatioti, A., Liakakou, E., Hatzianastassiou, N., Sciare, J., Mihalopoulos, N., and Bourtsoukidis, E.: Emission of volatile organic compounds from residential biomass burning and their rapid chemical transformations, *Science of The Total Environment*, 903, 166592, <https://doi.org/10.1016/j.scitotenv.2023.166592>, 2023.

955 Dimitriou, K. and Kassomenos, P.: Background concentrations of benzene, potential long range transport influences and corresponding cancer risk in four cities of central Europe, in relation to air mass origination, *J Environ Manage*, 262, 110374, <https://doi.org/10.1016/j.jenvman.2020.110374>, 2020.

960 Dorokhov, Y. L., Sheshukova, E. V., and Komarova, T. V.: Methanol in Plant Life, *Front Plant Sci*, 9, <https://doi.org/10.3389/fpls.2018.01623>, 2018.

Draxler, R. R. and H. G. D.: Description of the HYSPLIT_4 modeling system., Silver Spring, 24 pp., 1997.

965 Edtbauer, A., Stöner, C., Pfannerstill, E. Y., Berasategui, M., Walter, D., Crowley, J. N., Lelieveld, J., and Williams, J.: A new marine biogenic emission: methane sulfonamide (MSAM), dimethyl sulfide (DMS), and dimethyl sulfone (DMSO₂) measured in air over the Arabian Sea, *Atmos Chem Phys*, 20, 6081–6094, <https://doi.org/10.5194/acp-20-6081-2020>, 2020.

European Forest Fire Information System (EFFIS), European Commission, Copernicus Emergency Management Service: Annual fire reports, available at: <https://forest-fire.emergency.copernicus.eu/reports-and-publications/annual-fire-reports>, last access: 27

970 November 2025.

Ehn, M., Thornton, J. A., Kleist, E., Sipilä, M., Junninen, H., Pullinen, I., Springer, M., Rubach, F., Tillmann, R., Lee, B., Lopez-Hilfiker, F., Andres, S., Acir, I.-H., Rissanen, M., Jokinen, T., Schobesberger, S., Kangasluoma, J., Kontkanen, J., Nieminen, T., Kurtén, T., Nielsen, L. B., Jørgensen, S., Kjaergaard, H. G., Canagaratna, M., Maso, M. D., Berndt, T., Petäjä, T., Wahner, A.,

975 Kerminen, V.-M., Kulmala, M., Worsnop, D. R., Wildt, J., and Mentel, T. F.: A large source of low-volatility secondary organic aerosol, *Nature*, 506, 476–479, <https://doi.org/10.1038/nature13032>, 2014.

Emekwuru, N. and Ejohwomu, O.: Temperature, Humidity and Air Pollution Relationships during a Period of Rainy and Dry Seasons in Lagos, West Africa, *Climate*, 11, 113, <https://doi.org/10.3390/cli11050113>, 2023.

980 Fortunati, A., Barta, C., Brilli, F., Centritto, M., Zimmer, I., Schnitzler, J., and Loreto, F.: Isoprene emission is not temperature-dependent during and after severe drought-stress: a physiological and biochemical analysis, *The Plant Journal*, 55, 687–697, <https://doi.org/10.1111/j.1365-313X.2008.03538.x>, 2008.

985 Gelencsér, A., Siszler, K., and Hlavay, J.: Toluene–Benzene Concentration Ratio as a Tool for Characterizing the Distance from Vehicular Emission Sources, *Environ Sci Technol*, 31, 2869–2872, <https://doi.org/10.1021/es970004c>, 1997.

Georgiou, G. K., Kushta, J., Christoudias, T., Proestos, Y., and Lelieveld, J.: Air quality modelling over the Eastern Mediterranean: Seasonal sensitivity to anthropogenic emissions, *Atmos Environ*, 222, 117119, <https://doi.org/10.1016/j.atmosenv.2019.117119>, 2020.

990 Georgiou, G. K., Christoudias, T., Proestos, Y., Kushta, J., Pikridas, M., Sciare, J., Savvides, C., and Lelieveld, J.: Evaluation of WRF-Chem model (v3.9.1.1) real-time air quality forecasts over the Eastern Mediterranean, *Geosci Model Dev*, 15, 4129–4146, <https://doi.org/10.5194/gmd-15-4129-2022>, 2022.

995 Germain-Piaulenne, E., Paris, J.-D., Gros, V., Quéhé, P.-Y., Pikridas, M., Baisnée, D., Berchet, A., Sciare, J., and Bourtsoukidis, E.: Middle East oil and gas methane emissions signature captured at a remote site using light hydrocarbon tracers, *Atmos Environ X*, 22, 100253, <https://doi.org/10.1016/j.aeaoa.2024.100253>, 2024.

Gjstelund, I., Hollund, B. E., Kirkeleit, J., Daling, P. S., Sørheim, K. R., and Bråttveit, M.: Determinants of airborne benzene evaporating from fresh crude oils released into seawater, *Mar Pollut Bull*, 140, 395–402, <https://doi.org/10.1016/j.marpolbul.2018.12.045>, 2019.

1000 Gu, S., Guenther, A., and Faiola, C.: Effects of Anthropogenic and Biogenic Volatile Organic Compounds on Los Angeles Air Quality, *Environ Sci Technol*, 55, 12191–12201, <https://doi.org/10.1021/acs.est.1c01481>, 2021.

1005 Guo, H., Ling, Z. H., Simpson, I. J., Blake, D. R., and Wang, D. W.: Observations of isoprene, methacrolein (MAC) and methyl vinyl ketone (MVK) at a mountain site in Hong Kong, *Journal of Geophysical Research: Atmospheres*, 117, <https://doi.org/10.1029/2012JD017750>, 2012.

Holzinger, R., Williams, J., Salisbury, G., Klüpfel, T., de Reus, M., Traub, M., Crutzen, P. J., and Lelieveld, J.: Oxygenated compounds in aged biomass burning plumes over the Eastern

1010 Mediterranean: evidence for strong secondary production of methanol and acetone, *Atmos Chem Phys*, 5, 39–46, <https://doi.org/10.5194/acp-5-39-2005>, 2005.

Huang, X.-F., Zhang, B., Xia, S.-Y., Han, Y., Wang, C., Yu, G.-H., and Feng, N.: Sources of oxygenated volatile organic compounds (OVOCs) in urban atmospheres in North and South China, *Environmental Pollution*, 261, 114152, <https://doi.org/10.1016/j.envpol.2020.114152>, 2020.

1015 Huangfu, Y., Yuan, B., Wang, S., Wu, C., He, X., Qi, J., de Gouw, J., Warneke, C., Gilman, J. B., Wisthaler, A., Karl, T., Graus, M., Jobson, B. T., and Shao, M.: Revisiting Acetonitrile as Tracer of Biomass Burning in Anthropogenic-Influenced Environments, *Geophys Res Lett*, 48, <https://doi.org/10.1029/2020GL092322>, 2021.

- 1020 Kajos, M. K., Rantala, P., Hill, M., Hellén, H., Aalto, J., Patokoski, J., Taipale, R., Hoerger, C. C., Reimann, S., Ruuskanen, T. M., Rinne, J., and Petäjä, T.: Ambient measurements of aromatic and oxidized VOCs by PTR-MS and GC-MS: intercomparison between four instruments in a boreal forest in Finland, *Atmos Meas Tech*, 8, 4453–4473, <https://doi.org/10.5194/amt-8-4453-2015>, 2015.
- 1025 Kaltsonoudis, C., Kostenidou, E., Florou, K., Psichoudaki, M., and Pandis, S. N.: Temporal variability and sources of VOCs in urban areas of the eastern Mediterranean, *Atmos Chem Phys*, 16, 14825–14842, <https://doi.org/10.5194/acp-16-14825-2016>, 2016.
- Kamal, M. S., Razzak, S. A., and Hossain, M. M.: Catalytic oxidation of volatile organic compounds (VOCs) – A review, *Atmos Environ*, 140, 117–134, <https://doi.org/10.1016/j.atmosenv.2016.05.031>, 2016.
- 1030 Kiendler-Scharr, A., Wildt, J., Maso, M. D., Hohaus, T., Kleist, E., Mentel, T. F., Tillmann, R., Uerlings, R., Schurr, U., and Wahner, A.: New particle formation in forests inhibited by isoprene emissions, *Nature*, 461, 381–384, <https://doi.org/10.1038/nature08292>, 2009.
- Kleanthous, S., Vrekoussis, M., Mihalopoulos, N., Kalabokas, P., and Lelieveld, J.: On the temporal and spatial variation of ozone in Cyprus, *Science of The Total Environment*, 476–477, 677–687, <https://doi.org/10.1016/j.scitotenv.2013.12.101>, 2014.
- 1035 Knote, C., Hodzic, A., and Jimenez, J. L.: The effect of dry and wet deposition of condensable vapors on secondary organic aerosol concentrations over the continental United States, *Atmos. Chem. Phys.*, 15, 1–18, <https://doi.org/10.5194/acp-15-1-2015>, 2015a.
- 1040 Knote, C., Tuccella, P., Curci, G., Emmons, L., Orlando, J. J., Madronich, S., et al.: Influence of the choice of gas-phase mechanism on predictions of key gaseous pollutants during the AQMEII phase-2 intercomparison, *Atmos. Environ.*, 115, 553–568, <https://doi.org/10.1016/j.atmosenv.2014.11.066>, 2015b.
- Koppmann, R.: Chemistry of Volatile Organic Compounds in the Atmosphere, in: *Hydrocarbons, Oils and Lipids: Diversity, Origin, Chemistry and Fate*, Springer International Publishing, Cham, 811–822, https://doi.org/10.1007/978-3-319-90569-3_24, 2020.
- 1045 Koppmann, R., von Czapiewski, K., and Reid, J. S.: A review of biomass burning emissions, part I: gaseous emissions of carbon monoxide, methane, volatile organic compounds, and nitrogen containing compounds, <https://doi.org/10.5194/acpd-5-10455-2005>, 24 October 2005.
- 1050 Lazoglou, G., Hadjinicolaou, P., Sofokleous, I., Bruggeman, A., and Zittis, G.: Climate change and extremes in the Mediterranean island of Cyprus: from historical trends to future projections, *Environ Res Commun*, 6, 095020, <https://doi.org/10.1088/2515-7620/ad7927>, 2024.
- Lee, J. D., Squires, F. A., Sherwen, T., Wilde, S. E., Cliff, S. J., Carpenter, L. J., Hopkins, J. R., Bauguitte, S. J., Reed, C., Barker, P., Allen, G., Bannan, T. J., Matthews, E., Mehra, A., Percival, C., Heard, D. E., Whalley, L. K., Ronnie, G. V., Seldon, S., Ingham, T., Keller, C. A., Knowland, K. E., Nisbet, E. G., and Andrews, S.: Ozone production and precursor emission from wildfires in Africa, *Environmental Science: Atmospheres*, 1, 524–542, <https://doi.org/10.1039/D1EA00041A>, 2021.
- 1055 Lelieveld, J., Berresheim, H., Borrmann, S., Crutzen, P. J., Dentener, F. J., Fischer, H., Feichter, J., Flatau, P. J., Heland, J., Holzinger, R., Korrmann, R., Lawrence, M. G., Levin, Z., Markowicz, K. M., Mihalopoulos, N., Minikin, A., Ramanathan, V., de Reus, M., Roelofs, G. J., Scheeren, H. A., Sciare, J., Schlager, H., Schultz, M., Siegmund, P., Steil, B., Stephanou, E. G., Stier, P., Traub, M., Warneke, C., Williams, J., and Ziereis, H.: Global Air Pollution Crossroads over the Mediterranean, *Science* (1979), 298, 794–799, <https://doi.org/10.1126/science.1075457>, 2002.
- 1065 Liakakou, E., Vrekoussis, M., Bonsang, B., Donousis, Ch., Kanakidou, M., and Mihalopoulos, N.: Isoprene above the Eastern Mediterranean: Seasonal variation and contribution to the oxidation capacity of the atmosphere, *Atmos Environ*, 41, 1002–1010, <https://doi.org/10.1016/j.atmosenv.2006.09.034>, 2007.
- Liakakou, E., Bonsang, B., Williams, J., Kalivitis, N., Kanakidou, M., and Mihalopoulos, N.: C2–C8 NMHCs over the Eastern Mediterranean: Seasonal variation and impact on regional oxidation chemistry, *Atmos Environ*, 43, 5611–5621, <https://doi.org/10.1016/j.atmosenv.2009.07.067>, 2009.
- 1070 Liu, Y., Shao, M., Kuster, W. C., Goldan, P. D., Li, X., Lu, S., and Gouw, J. A. de: Source Identification of Reactive Hydrocarbons and Oxygenated VOCs in the Summertime in Beijing, *Environ Sci Technol*, 43, 75–81, <https://doi.org/10.1021/es801716n>, 2009.
- 1075 Matthew, O. J.: Estimation of diurnal patterns of global solar radiation, temperature, relative humidity and wind speed from daily datasets at a humid tropical location, *Agric For Meteorol*, 322, 109003, <https://doi.org/10.1016/j.agrformet.2022.109003>, 2022.
- Mecca, M., Todaro, L., D’Auria, M., Italiano, S. S. P., Sofo, A., and Ripullone, F.: Volatile Organic Compounds (VOCs) in Mediterranean Oak Forests of Hungarian Oak (*Quercus frainetto* Ten) Affected by Dieback Phenomena, *Forests*, 15, 1072, <https://doi.org/10.3390/f15061072>, 2024.
- 1080

- Mellouki, A., Wallington, T. J., and Chen, J.: Atmospheric Chemistry of Oxygenated Volatile Organic Compounds: Impacts on Air Quality and Climate, *Chem Rev*, 115, 3984–4014, <https://doi.org/10.1021/cr500549n>, 2015.
- 1085 Misztal, P. K., Hewitt, C. N., Wildt, J., Blande, J. D., Eller, A. S. D., Fares, S., Gentner, D. R., Gilman, J. B., Graus, M., Greenberg, J., Guenther, A. B., Hansel, A., Harley, P., Huang, M., Jardine, K., Karl, T., Kaser, L., Keutsch, F. N., Kiendler-Scharr, A., Kleist, E., Lerner, B. M., Li, T., Mak, J., Nölscher, A. C., Schnitzhofer, R., Sinha, V., Thornton, B., Warneke, C., Wegener, F., Werner, C., Williams, J., Worton, D. R., Yassaa, N., and Goldstein, A. H.: Atmospheric benzenoid emissions from plants rival those from fossil fuels, *Sci Rep*, 5, 12064, <https://doi.org/10.1038/srep12064>, 2015.
- 1090 Monson, R. K., Jaeger, C. H., Adams, W. W., Driggers, E. M., Silver, G. M., and Fall, R.: Relationships among Isoprene Emission Rate, Photosynthesis, and Isoprene Synthase Activity as Influenced by Temperature, *Plant Physiol*, 98, 1175–1180, <https://doi.org/10.1104/pp.98.3.1175>, 1992.
- 1095 Mukherjee, S., Pandithurai, G., Waghmare, V., Mahajan, A. S., Tinel, L., Aslam, M. Y., Meena, G. S., Patil, S., Buchunde, P., and Kumar, A.: Seasonal variability of volatile organic compounds (VOCs) at a high-altitude station in the Western Ghats, India: Influence of biogenic, anthropogenic emissions and long-range transport, *Atmos Environ*, 331, 120598, <https://doi.org/10.1016/j.atmosenv.2024.120598>, 2024.
- 1100 Pagonis, D., Sekimoto, K., and de Gouw, J.: A Library of Proton-Transfer Reactions of H_3O^+ Ions Used for Trace Gas Detection, *J Am Soc Mass Spectrom*, 30, 1330–1335, <https://doi.org/10.1007/s13361-019-02209-3>, 2019.
- 1105 Panopoulou, A., Liakakou, E., Gros, V., Sauvage, S., Locoge, N., Bonsang, B., Psiloglou, B. E., Gerasopoulos, E., and Mihalopoulos, N.: Non-methane hydrocarbon variability in Athens during wintertime: the role of traffic and heating, *Atmos Chem Phys*, 18, 16139–16154, <https://doi.org/10.5194/acp-18-16139-2018>, 2018.
- 1110 Panopoulou, A., Liakakou, E., Sauvage, S., Gros, V., Locoge, N., Stavroulas, I., Bonsang, B., Gerasopoulos, E., and Mihalopoulos, N.: Yearlong measurements of monoterpenes and isoprene in a Mediterranean city (Athens): Natural vs anthropogenic origin, *Atmos Environ*, 243, 117803, <https://doi.org/10.1016/j.atmosenv.2020.117803>, 2020.
- Paton-Walsh, C., Wilson, S. R., Jones, N. B., and Griffith, D. W. T.: Measurement of methanol emissions from Australian wildfires by ground-based solar Fourier transform spectroscopy, *Geophys Res Lett*, 35, <https://doi.org/10.1029/2007GL032951>, 2008.
- 1115 Pennington, E. A., Seltzer, K. M., Murphy, B. N., Qin, M., Seinfeld, J. H., and Pye, H. O. T.: Modeling secondary organic aerosol formation from volatile chemical products, *Atmos Chem Phys*, 21, 18247–18261, <https://doi.org/10.5194/acp-21-18247-2021>, 2021.
- Peñuelas, J. and Staudt, M.: BVOCs and global change, *Trends Plant Sci*, 15, 133–144, <https://doi.org/10.1016/j.tplants.2009.12.005>, 2010.
- 1120 Pérez, I. A., García, M. Á., Sánchez, M. L., Pardo, N., and Fernández-Duque, B.: Key Points in Air Pollution Meteorology, *Int J Environ Res Public Health*, 17, 8349, <https://doi.org/10.3390/ijerph17228349>, 2020.
- Perrone, M. G., Carbone, C., Faedo, D., Ferrero, L., Maggioni, A., Sangiorgi, G., and Bolzacchini, E.: Exhaust emissions of polycyclic aromatic hydrocarbons, n-alkanes and phenols from vehicles coming within different European classes, *Atmos Environ*, 82, 391–400, <https://doi.org/10.1016/j.atmosenv.2013.10.040>, 2014.
- 1125 Pikridas, M., Vrekoussis, M., Sciare, J., Kleanthous, S., Vasiliadou, E., Kizas, C., Savvides, C., and Mihalopoulos, N.: Spatial and temporal (short and long-term) variability of submicron, fine and sub-10 μm particulate matter (PM₁, PM_{2.5}, PM₁₀) in Cyprus, *Atmos Environ*, 191, 79–93, <https://doi.org/10.1016/j.atmosenv.2018.07.048>, 2018.
- 1130 Pinthong, N., Thepanondh, S., and Kondo, A.: Source Identification of VOCs and their Environmental Health Risk in a Petrochemical Industrial Area, *Aerosol Air Qual Res*, 22, 210064, <https://doi.org/10.4209/aaqr.210064>, 2022.
- Pochanart, P., Akimoto, H., Kajii, Y., Potemkin, V. M., and Khodzher, T. V.: Regional background ozone and carbon monoxide variations in remote Siberia/East Asia, *Journal of Geophysical Research: Atmospheres*, 108, <https://doi.org/10.1029/2001JD001412>, 2003.
- 1135 Pugliese, G., Ingrischi, J., Meredith, L. K., Pfannerstill, E. Y., Klüpfel, T., Meeran, K., Byron, J., Purser, G., Gil-Loaiza, J., van Haren, J., Dontsova, K., Kreuzwieser, J., Ladd, S. N., Werner, C., and Williams, J.: Effects of drought and recovery on soil volatile organic compound fluxes in an experimental rainforest, *Nat Commun*, 14, 5064, <https://doi.org/10.1038/s41467-023-40661-8>, 2023.
- 1140

- Ralf Kopppmann: Volatile Organic Compounds in the Atmosphere, edited by: Kopppmann, R., Wiley, <https://doi.org/10.1002/9780470988657>, 2007.
- Robles, H.: Acetonitrile, in: Encyclopedia of Toxicology, Elsevier, 28–30, <https://doi.org/10.1016/B0-12-369400-0/00015-6>, 2005.
- 1145 Salisbury, G., Williams, J., Holzinger, R., Gros, V., Mihalopoulos, N., Vrekoussis, M., Sarda-Estève, R., Berresheim, H., von Kuhlmann, R., Lawrence, M., and Lelieveld, J.: Ground-based PTR-MS measurements of reactive organic compounds during the MINOS campaign in Crete, July–August 2001, *Atmos Chem Phys*, 3, 925–940, <https://doi.org/10.5194/acp-3-925-2003>, 2003.
- 1150 Schieweck, A., Uhde, E., and Salthammer, T.: Determination of acrolein in ambient air and in the atmosphere of environmental test chambers, *Environ Sci Process Impacts*, 23, 1729–1746, <https://doi.org/10.1039/D1EM00221J>, 2021.
- Sciare J: The Agia Marina Xyliatou Observatory: A remote supersite in Cyprus to monitor changes in the atmospheric composition of the Eastern Mediterranean and the Middle East, in: EGU General Assembly Conference Abstracts, 2016.
- 1155 Sicre, M. A., Marty, J. C., Salot, A., Aparicio, X., Grimalt, J., and Albaiges, J.: Aliphatic and Aromatic Hydrocarbons in the Mediterranean Aerosol, *Int J Environ Anal Chem*, 29, 73–94, <https://doi.org/10.1080/03067318708078412>, 1987.
- Sindelarova, K., Granier, C., Bouarar, I., Guenther, A., Tilmes, S., Stavrakou, T., Müller, J.-F., Kuhn, U., Stefani, P., and Knorr, W.: Global data set of biogenic VOC emissions calculated by the MEGAN model over the last 30 years, *Atmos Chem Phys*, 14, 9317–9341, <https://doi.org/10.5194/acp-14-9317-2014>, 2014.
- 1160 Song, W., Williams, J., Yassaa, N., Martinez, M., Carnero, J. A. A., Hidalgo, P. J., Bozem, H., and Lelieveld, J.: Winter and summer characterization of biogenic enantiomeric monoterpenes and anthropogenic BTEX compounds at a Mediterranean Stone Pine forest site, *J Atmos Chem*, 68, 233–250, <https://doi.org/10.1007/s10874-012-9219-4>, 2011.
- 1165 Soukissian, T. and Sotiriou, M.-A.: Long-Term Variability of Wind Speed and Direction in the Mediterranean Basin, *Wind*, 2, 513–534, <https://doi.org/10.3390/wind2030028>, 2022.
- Stein, A. F., Draxler, R. R., Rolph, G. D., Stunder, B. J. B., Cohen, M. D., and Ngan, F.: NOAA's HYSPLIT Atmospheric Transport and Dispersion Modeling System, *Bull Am Meteorol Soc*, 96, 2059–2077, <https://doi.org/10.1175/BAMS-D-14-00110.1>, 2015.
- 1170 Stockwell, W. R., Kirchner, F., Kuhn, M., and Seefeld, S.: A new mechanism for regional atmospheric chemistry modeling, *Journal of Geophysical Research: Atmospheres*, 102, 25847–25879, <https://doi.org/10.1029/97JD00849>, 1997.
- 1175 Strada, S., Pozzer, A., Giuliani, G., Coppola, E., Solmon, F., Jiang, X., Guenther, A., Bourtsoukidis, E., Serça, D., Williams, J., and Giorgi, F.: Assessment of isoprene and near-surface ozone sensitivities to water stress over the Euro-Mediterranean region, *Atmos Chem Phys*, 23, 13301–13327, <https://doi.org/10.5194/acp-23-13301-2023>, 2023.
- Thunis, P., Crippa, M., Cuvelier, C., Guizzardi, D., de Meij, A., Oreggioni, G., and Pisoni, E.: Sensitivity of air quality modelling to different emission inventories: a case study over Europe, *Atmos. Environ. X*, 10, 100111, 2021.
- 1180 Tiwari, V., Hanai, Y., and Masunaga, S.: Ambient levels of volatile organic compounds in the vicinity of petrochemical industrial area of Yokohama, Japan, *Air Qual Atmos Health*, 3, 65–75, <https://doi.org/10.1007/s11869-009-0052-0>, 2010.
- 1185 Torres-Vinces, L., Contreras-Zarazua, G., Huerta-Rosas, B., Sánchez-Ramírez, E., and Segovia-Hernández, J. G.: Methyl Ethyl Ketone Production through an Intensified Process, *Chem Eng Technol*, 43, 1433–1441, <https://doi.org/10.1002/ceat.201900664>, 2020.
- Traub, M., Fischer, H., de Reus, M., Kormann, R., Heland, H., Ziereis, H., Schlager, H., Holzinger, R., Williams, J., Warneke, C., de Gouw, J., and Lelieveld, J.: Chemical characteristics assigned to trajectory clusters during the MINOS campaign, *Atmos Chem Phys*, 3, 459–468, <https://doi.org/10.5194/acp-3-459-2003>, 2003.
- 1190 Vlasenko, A., Macdonald, A. . M., Sjostedt, S. J., and Abbatt, J. P. D.: Formaldehyde measurements by Proton transfer reaction – Mass Spectrometry (PTR-MS): correction for humidity effects, *Atmos Meas Tech*, 3, 1055–1062, <https://doi.org/10.5194/amt-3-1055-2010>, 2010.
- 1195 Vrekoussis, M., Pikridas, M., Rousogenous, C., Christodoulou, A., Desservettaz, M., Sciare, J., Richter, A., Bougoudis, I., Savvides, C., and Papadopoulos, C.: Local and regional air pollution characteristics in Cyprus: A long-term trace gases observations analysis, *Science of The Total Environment*, 845, 157315, <https://doi.org/10.1016/j.scitotenv.2022.157315>, 2022.
- 1200 Wang, H., Ma, X., Tan, Z., Wang, H., Chen, X., Chen, S., Gao, Y., Liu, Y., Liu, Y., Yang, X., Yuan, B., Zeng, L., Huang, C., Lu, K., and Zhang, Y.: Anthropogenic monoterpenes aggravating ozone pollution, *Natl Sci Rev*, 9, <https://doi.org/10.1093/nsr/nwac103>, 2022a.

- Wang, L., Lun, X., Wang, Q., and Wu, J.: Biogenic volatile organic compounds emissions, atmospheric chemistry, and environmental implications: a review, *Environ Chem Lett*, 22, 3033–3058, <https://doi.org/10.1007/s10311-024-01785-5>, 2024.
- 1205 Wang, N., Edtbauer, A., Stönnner, C., Pozzer, A., Bourtsoukidis, E., Ernle, L., Dienhart, D., Hottmann, B., Fischer, H., Schuladen, J., Crowley, J. N., Paris, J.-D., Lelieveld, J., and Williams, J.: Measurements of carbonyl compounds around the Arabian Peninsula: overview and model comparison, *Atmos Chem Phys*, 20, 10807–10829, <https://doi.org/10.5194/acp-20-10807-2020>, 2020.
- 1210 Wang, S., Yuan, B., Wu, C., Wang, C., Li, T., He, X., Huangfu, Y., Qi, J., Li, X.-B., Sha, Q., Zhu, M., Lou, S., Wang, H., Karl, T., Graus, M., Yuan, Z., and Shao, M.: Oxygenated volatile organic compounds (VOCs) as significant but varied contributors to VOC emissions from vehicles, *Atmos Chem Phys*, 22, 9703–9720, <https://doi.org/10.5194/acp-22-9703-2022>, 2022b.
- 1215 Wang, H., Lu, X., Seco, R., Stavrakou, T., Karl, T., Jiang, X., et al.: Modeling isoprene emission response to drought and heatwaves within MEGAN using evapotranspiration data and by coupling with the Community Land Model, *J. Adv. Model. Earth Syst.*, **14**, e2022MS003174, <https://doi.org/10.1029/2022MS003174>, 2022c.
- 1220 Xu, Z., Zou, Q., Jin, L., Shen, Y., Shen, J., Xu, B., Qu, F., Zhang, F., Xu, J., Pei, X., Xie, G., Kuang, B., Huang, X., Tian, X., and Wang, Z.: Characteristics and sources of ambient Volatile Organic Compounds (VOCs) at a regional background site, YRD region, China: Significant influence of solvent evaporation during hot months, *Science of The Total Environment*, 857, 159674, <https://doi.org/10.1016/j.scitotenv.2022.159674>, 2023.
- 1225 Yáñez-Serrano, A. M., Nölscher, A. C., Bourtsoukidis, E., Derstroff, B., Zannoni, N., Gros, V., Lanza, M., Brito, J., Noe, S. M., House, E., Hewitt, C. N., Langford, B., Nemitz, E., Behrendt, T., Williams, J., Artaxo, P., Andreae, M. O., and Kesselmeier, J.: Atmospheric mixing ratios of methyl ethyl ketone (2-butanone) in tropical, boreal, temperate and marine environments, *Atmos Chem Phys*, 16, 10965–10984, <https://doi.org/10.5194/acp-16-10965-2016>, 2016.
- 1230 Yáñez-Serrano, A. M., Filella, I., LLusà, J., Gargallo-Garriga, A., Granda, V., Bourtsoukidis, E., Williams, J., Seco, R., Cappellin, L., Werner, C., de Gouw, J., and Peñuelas, J.: GLOVOCS - Master compound assignment guide for proton transfer reaction mass spectrometry users, *Atmos Environ*, 244, 117929, <https://doi.org/10.1016/j.atmosenv.2020.117929>, 2021.
- 1235 Yokelson, R. J., Goode, J. G., Ward, D. E., Susott, R. A., Babbitt, R. E., Wade, D. D., Bertschi, I., Griffith, D. W. T., and Hao, W. M.: Emissions of formaldehyde, acetic acid, methanol, and other trace gases from biomass fires in North Carolina measured by airborne Fourier transform infrared spectroscopy, *Journal of Geophysical Research: Atmospheres*, 104, 30109–30125, <https://doi.org/10.1029/1999JD900817>, 1999.
- 1240 Yuan, Q., Zhang, Z., Chen, Y., Hui, L., Wang, M., Xia, M., Zou, Z., Wei, W., Ho, K. F., Wang, Z., Lai, S., Zhang, Y., Wang, T., and Lee, S.: Origin and transformation of volatile organic compounds at a regional background site in Hong Kong: Varied photochemical processes from different source regions, *Science of The Total Environment*, 908, 168316, <https://doi.org/10.1016/j.scitotenv.2023.168316>, 2024.
- 1245 Zittis, G., Almazroui, M., Alpert, P., Ciais, P., Cramer, W., Dahdal, Y., Fnais, M., Francis, D., Hadjinicolaou, P., Howari, F., Jrrar, A., Kaskaoutis, D. G., Kulmala, M., Lazoglou, G., Mihalopoulos, N., Lin, X., Rudich, Y., Sciare, J., Stenchikov, G., Xoplaki, E., and Lelieveld, J.: Climate Change and Weather Extremes in the Eastern Mediterranean and Middle East, *Reviews of Geophysics*, 60, <https://doi.org/10.1029/2021RG000762>, 2022.