

# Observations of nanoparticle shrinkage phenomena

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**Abstract.** Atmospheric new particle formation (NPF) is a major source of aerosol particles in the Earth's atmosphere. However, process-level understanding of the early stages of particle formation and growth remains poorly represented in climate models, limiting accurate estimates of aerosol effective radiative forcing. Here, we use comprehensive observations from the Spring Particles in Cyprus (SPICY) field campaign conducted at a rural background site in Cyprus. We report new observations of nanoparticle shrinkage (NPS), marked by the rapid decrease in size of sub-20 nm particles occurring in the absence of preceding NPF event. Thus, the particle size distributions exhibit a mirror image of the conventional "banana-shaped" NPF pattern, forming a distinctive "reverse-NPF" pattern. We identified three NPS events during the campaign and show that this phenomenon is not primarily driven by low concentrations of condensable vapours, their scavenging by pre-existing particles, or primary nanoparticle sources. Instead, it is associated with atmospheric dilution, as indicated by air-mass trajectory analysis. Fast-moving air masses can enhance turbulent mixing, leading to dilution of particle number concentrations and changes in particle size distributions. Together with volatility-resolved analysis, these results suggest that NPS is governed by atmospheric dilution, which reduces particle-phase organic mass and shift the gas-particle equilibrium toward evaporation, with contributions dominated by organic compounds of low and moderate volatility. Our results demonstrate that NPS events provide a previously unrecognised sink for nanoparticles, which are controlled by air-mass dynamics and organic vapour volatility.

## 1. Introduction

Atmospheric aerosols remain one of the largest uncertainties in quantifying effective radiative forcing and, consequently, predictions of climate change. Aerosols affect climate directly by scattering and absorbing solar radiation and indirectly by serving as cloud condensation nuclei (CCN), thereby influencing cloud properties and

precipitation patterns (Twomey, 1977; Charlson et al., 1992; Rosenfeld et al., 2014; Ipcc, 2023). Atmospheric  
40 new particle formation (NPF) from low-volatility condensable vapours is prevalent throughout the troposphere  
(Kulmala et al., 2000; Zhang et al., 2004; Clarke, 1993), and represents the dominant global source of aerosol  
number concentrations. NPF occurs under a broad spectrum of atmospheric environments, from urban and  
forested sites to marine and free-tropospheric conditions (Kerminen et al., 2018; Kulmala et al., 2004; Zhao et al.,  
2024). Atmospheric observations indicate that 10-60% of NPF events contribute to CCN formation, enhancing  
45 their concentrations by factors that range from 0.5 to 11 at regional scales (Kerminen et al., 2018; Laakso et al.,  
2013; Asmi et al., 2011; Sebastian et al., 2022; Sihto et al., 2011; Rose et al., 2017; Williamson et al., 2019).  
Global model simulations also suggest that NPF accounts for roughly half of the tropospheric CCN population  
(Gordon et al., 2017; Merikanto et al., 2009; Yu and Luo, 2009). NPF events and nanoparticle growth have also  
been linked to severe haze episodes that exacerbate air pollution (Kulmala et al., 2022; Mishra et al., 2023; Guo  
50 et al., 2014). However, limited understanding of which condensable vapours dominate the phenomena, the initial  
stages of particle formation and growth, and the survival probability of nanoparticles hinder accurate  
representation of particle evolution from a few nanometers to CCN-active sizes (>50-100 nm) in models.

Decreasing mode diameter (DMD; also referred to as particle shrinkage) events are characterised by an evolution  
55 of particle size distributions during NPF, in which the modal diameter of nucleation-mode particles initially  
increases into the Aitken-mode size range and subsequently decreases to smaller sizes. This results in an inverted  
U-shaped temporal evolution of the modal diameter (Kamra et al., 2022; Hakala et al., 2019; Alonso-Blanco et  
al., 2017; Salma et al., 2016; Skrabalova et al., 2015; Cusack et al., 2013; Young et al., 2013; Yao et al., 2010;  
Backman et al., 2012; Hussein et al., 2020; Zhang et al., 2016; Yue et al., 2016; Kivekäs et al., 2016). As a result,  
60 DMD events can substantially suppress CCN production by preventing particles from reaching activation sizes.  
Hakala et al. (2023) attributed DMD events to low-growth environments associated with reduced photochemical  
productions of condensable vapours. Limited or slow particle growth rate, together with surface tension effects,  
can inhibit CCN activation during NPF events (Cai et al., 2021), a process that remains underrepresented in current  
atmospheric process-based models.

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Observations across diverse regions reveal that DMD events arise from distinct mechanisms influenced by local-  
to-regional conditions and photochemical processes. Observations in East Asia (Hong Kong and Taiwan) suggest  
that DMD events are often driven by the evaporation of condensed semi-volatile compounds under varying  
photochemical conditions (Yao et al., 2010; Young et al., 2013), while studies in European urban areas (Budapest,  
70 Prague, and Madrid) and densely forested environments in the western Mediterranean (Montseny) attribute these  
events to a combination of late-afternoon reductions in condensable vapours, diminished photochemistry,  
evaporation of semi-volatile compounds, and increased atmospheric dilution (Salma et al., 2016; Alonso-Blanco  
et al., 2017; Skrabalova et al., 2015; Cusack et al., 2013). In cleaner environments, such as the city of Toronto,  
limited availability of low-volatility vapours was found to inhibit particle growth, favoring DMD events compared  
75 to polluted locations like Qingdao (Zhu et al., 2014). El-Sayed et al. (2018) suggest that isoprene oxidation  
products under low oxides of nitrogen ( $\text{NO}_x$ ) conditions can reversibly partition into condensed water in the  
particle phase, forming transient secondary organic aerosol (SOA) prone to evaporation. At an urban background  
site in Amman, Jordan, DMD events were observed on nearly 50% of NPF days and were linked to variations in

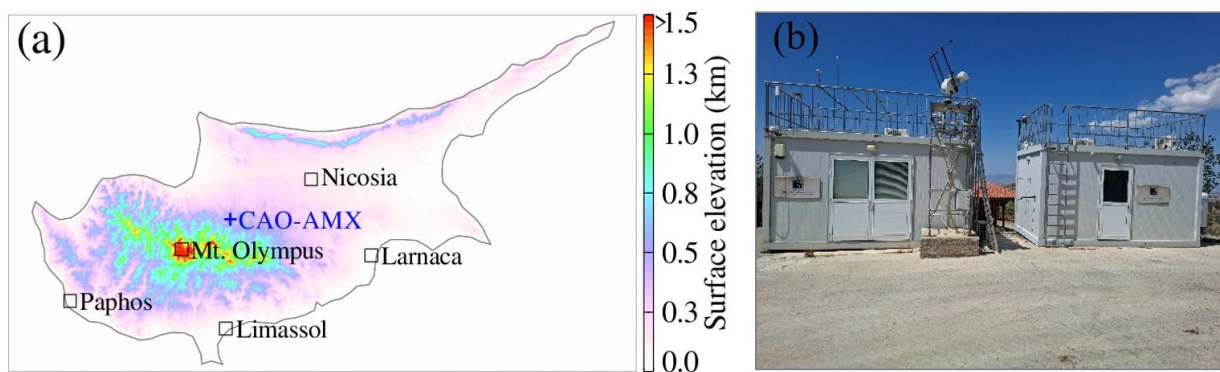
80 particle formation and growth rates during atmospheric transport (Hussein et al., 2020). At a rural arid site in  
Western Saudi Arabia (Hada Al Sham), DMD events were even more frequent, occurring on 76% of NPF days  
(Hakala et al., 2019). Such events are driven by changes in air-mass origin (Kivekäs et al., 2016), with less-grown  
particles arriving at the site after larger ones, while evaporation plays an insignificant role (Hakala et al., 2023).  
These observations indicate that DMD events result from a complex coupling of photochemical activity,  
condensing vapour availability, meteorological conditions, and air-mass history.

85 Here, we elucidate a new class of DMD events, which we refer to as nanoparticle shrinkage (NPS) events,  
characterised by a rapid decrease in particle mode diameter within the sub-20 nm size range that occurs in the  
absence of preceding NPF typically associated with DMD events. As a result, the particle size distributions display  
a mirror image of the conventional “banana-shaped” NPF pattern, forming a distinctive “reverse-NPF” signature.  
90 To our knowledge, this behaviour has not previously been examined in literature. We investigate these events  
using comprehensive measurements carried out during the Spring Particles In CYprus (SPICY) field campaign at  
the rural background site, Cyprus Atmospheric Observatory - Agia Marina Xyliatou (CAO-AMX) in spring 2024.

## 2. Methods

### 95 2.1 Measurement Site

Cyprus is the third most populous island in the Mediterranean. The SPICY field campaign was conducted from 1  
April to 15 June 2024 at the rural background site CAO-AMX (35.038692° N, 33.057850° E, 532 m a.m.s.l., Fig.  
1), with a focus on processes linking direct emissions to CCN formation in an environment where NPF events  
dominate (annual median > 58%) (Deot et al., 2025; Baalbaki et al., 2021). The campaign included an extensive  
100 suite of instruments for measuring condensing vapours, trace gases, VOCs, as well as particle size distributions  
and CCN concentrations. The CAO-AMX site is integrated into several international monitoring networks,  
including European Monitoring and Evaluation Programme (EMEP, Tørseth et al., 2012), Global Atmosphere  
Watch (GAW, Schultz et al., 2015), Aerosol, Clouds, and Trace Gases Research Infrastructure (ACTRIS, Laj et  
al., 2024), AEROSOL RObotic NETwork (AERONET, Holben et al., 1998), and EUMETNET Profiling Program  
105 (E-PROFILE, Illingworth et al., 2019), and is collocated with one of the nine stations of the Cyprus air quality  
monitoring network operated by the Department of Labour Inspection (DLI). Further details of CAO-AMX and  
other CAO network sites are available in Jokinen et al. (2026). The CAO-AMX site is characterised by a  
Mediterranean climate, with hot, dry summers and mild, wet winters. In spring, the median (range) values of  
temperature, relative humidity, solar radiation, and rainfall are 17.8 °C (2.4 - 37.6 °C), 51.2 % (7.6 - 99.2 %),  
110 448.4 W m<sup>-2</sup> (10.5 - 1027.4 W m<sup>-2</sup>), and 3.1 mm (1.2 - 24.6 mm), respectively (Jokinen et al., 2026). Anthropogenic  
emissions around the CAO-AMX site are minimal, with the nearest major cities located ~35 km northeast  
(Nicosia) and ~50 km southeast (Larnaca) of the site.



**Figure 1:** (a) Surface elevation map of Cyprus, showing the location of the CAO-AMX measurement station (blue marker) in the foothills of the Troodos Mountains. Major cities (Paphos, Limassol, Larnaca, and Nicosia) and Mount Olympus (the highest point in Cyprus) are indicated by open square boxes. Elevation data are taken from the U.S. Geological Survey global digital elevation model (DEM) with a horizontal resolution of 30 arc seconds (approximately 1 km; GTOPO30). (b) Image of the CAO-AMX station, consisting of two containers housing advanced instrumentation for condensing vapours, trace gases, and aerosols.

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## 2.2 Instrumentation and ancillary data

### 2.2.1. Multi-scheme chemical IONisation Atmospheric Pressure interface Time-Of-Flight mass spectrometer (MION-APi-TOF)

The APi-TOF (ToFwerk AG, Switzerland) is a high-resolution mass spectrometer capable of identifying ambient atmospheric ions and naturally occurring ion clusters with high mass accuracy and resolving power. The APi-TOF was equipped with a second-generation MION-2 inlet (Karsa Ltd., Finland), which allows rapid switching of reagent ions based on the chemical ionisation (CI)-inlet principle and ambient naturally charged ions (Rissanen et al., 2019). In this campaign, the MION-2 inlet was operated exclusively in negative polarity and operated with a 15-minute temporal resolution, switching between ambient ion mode for anion measurements and nitrate-based CI-mode for neutral compounds. The MION-APi-TOF sampled ambient air at 0.8 liter per minute (lpm) through a critical orifice (0.3 mm diameter). The sampled air was ionised in the MION source, and subsequently the gas-phase ions were transferred to the TOF chamber, where the charged compounds were separated according to their mass-to-charge ratio ( $m/z$ ) with a resolving power of about 9000 Th/Th. Data analysis was performed using the MATLAB-based *tofTools* package developed by Junninen et al. (2010). The  $C_{\text{sat}}$  values for individual HOMs were calculated based on their molecular composition using the parametrisation by Mohr et al. (2019).

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### 2.2.2 Proton Transfer Reaction Time-of-Flight mass spectrometer (PTR-ToF-MS)

Online measurements of VOCs were conducted using a high-resolution PTR-ToF-MS 4000 (Ionicon Analytik GmbH, Austria) following the methodology described by Desservettaz et al. (2023). The drift tube was operated at  $E/N = 128$  Td, 2.2 mbar, 630 V, and 85 °C. Ambient air was sampled at 0.21 lpm through a heated 2-m long PEEK line with a PTFE filter, minimizing losses by using inert materials (PTFE, PEEK, glass). Mass spectra was recorded every 30 s and averaged to 10 minutes. Background signals were acquired every two hours with VOC-free air from a platinum catalyst, and calibrations were performed bi-weekly with certified standards (Apel-Riemer Environmental, USA). Data were processed with the Ionicon Data Analyzer (IDA) and matched against

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145 the GLOVOCS database (Yáñez-Serrano et al., 2016), yielding ~76 reliably quantified VOCs after applying  
detection limits, calibration corrections, and quality filtering. Formaldehyde was excluded due to humidity-related  
detection artefacts(Vlasenko et al., 2010).

### **2.2.3 Nano condensation nucleus counter (nCNC)**

150 The A11 nCNC (Airmodus Oy, Finland) is composed of a particle size magnifier (PSM; Model A10) coupled  
with a CPC (Model A20) (Vanhanen et al., 2011). The sample inlet tube was approximately 60 cm long. The PSM  
was operated in scanning saturator flow mode (0.1 – 1.3 lpm), corresponding to a cutoff diameter of ~1.1 - 2.5  
155 nm. To minimize artefacts, the setup included an inlet system performing background (zero) checks three times  
daily (12 minutes each, equivalent to three full scans) and a core sampling piece to reduce line losses of sub-3 nm  
particles (Baalbaki et al., 2021). A dilution with dry particle-free air behind the inlet head was employed to ensure  
the dew point temperatures < 20 °C, following ACTRIS standard procedures for in-situ aerosol sampling and  
analyses (Lehtipalo et al., 2022).

### **2.2.4 Neutral Cluster and Air Ion Spectrometer (NAIS)**

160 Ion and particle size distributions were measured using a NAIS (Airel Ltd., Estonia). This instrument measures  
ion and naturally charged particle size distributions in the diameter range of 0.8–42 nm (109 size bins obtained  
from 25 electrometers) (Mirme and Mirme, 2013; Manninen et al., 2016), with measurements performed in both  
positive and negative polarities. The total particle size distribution was measured by charging aerosols using a  
corona charger, followed by size separation and detection carried out by two multichannel electrical mobility  
165 analyzer columns operated in parallel. Concentrations of particles with sizes smaller than ~2 nm were excluded  
due to known artefacts from the corona charger. Raw counts were inverted using the NAIS SPECTOPS algorithm  
and corrected for inlet losses (Gormley and Kennedy, 1948). Further details of the NAIS setup at CAO-AMX are  
available in *Deot et al. (2025)*.

### **2.2.5 Scanning Mobility Particle Sizer (SMPS)**

170 The SMPS is composed of a TSI 3083 long differential mobility analyzer (DMA) and a TSI 3789 CPC. It measures  
particle size distribution between 10 and 777 nm (61 size bins). The aerosol and sheath flows were checked weekly  
and set to 1.0 and 4.8 lpm, respectively. The SMPS sampled ambient air using a 3-m long sample inlet tube.  
Aerosol sample drying was achieved using a Nafion dryer (typically RH < 40%), following the ACTRIS standard  
175 procedures for in situ aerosol sampling and analyses, and charge neutralization was achieved using a TSI  
radioactive (<sup>85</sup>Kr) neutralizer (model 3077). The raw data from the SMPS was inverted using TSI's Aerosol  
Instrument Manager (AIM, version 11.5) software. The nano-DMA SMPS, consisting of a TSI 3085 DMA and a  
TSI water-based 3786 CPC, was also operated simultaneously, with a sample flow rate of 0.3 lpm. The nano-  
DMA SMPS sampled ambient air using a 2-m long sample inlet tube. It measures particle size distribution between  
180 3.1 and 105.5 nm (99 size bins). Additionally, a TSI butanol-based standalone 3750 CPC (D50 = 7 nm) was also  
operated, with a sample flow rate of 1 lpm.

### **2.2.6 Aethalometer**

BC mass concentrations were measured using a seven-wavelength aethalometer (AE33, Magee Scientific, USA) at 1-minute resolution. Ambient aerosol was sampled through a PM<sub>1</sub> inlet at 5 lpm after drying with a Nafion dryer. BC concentrations were derived at 880 nm using a mass absorption cross-section of 7.77 m<sup>2</sup> g<sup>-1</sup> (Petzold et al., 2013). This wavelength was chosen for calculating BC concentration because absorption by other aerosols is negligible at this wavelength (Drinovec et al., 2015). The aethalometer uses Teflon-coated glass fiber tape, and aerosols are collected onto two parallel spots for optical absorption measurements, which provide real-time compensation for the filter loading effect.

### ***2.2.7 Ceilometer***

The Vaisala Ceilometer CL51, in operation since 2021 and part of the E-PROFILE network, provides vertical profiles of aerosols and clouds (Münkel and Roininen, 2010). It employs an eye-safe InGaAs diode-laser lidar (910 ± 10 nm, 110 ns pulses, 6.5 kHz) in a vertical or near-vertical direction, detecting aerosols and clouds from ~300 m to 15 km with 10 m resolution. Backscatter profiles are processed with the BL-VIEW software, which applies a gradient method with a “cloud and precipitation filter” to remove spurious signals (Münkel and Roininen, 2010; Emeis et al., 2007). The software combines gradient and idealized backscatter approaches to automatically estimate planetary boundary layer (PBL) height at 16s temporal and 10 m vertical resolution. In this study, we retained Level 3 boundary layer height data with a “good” quality control index.

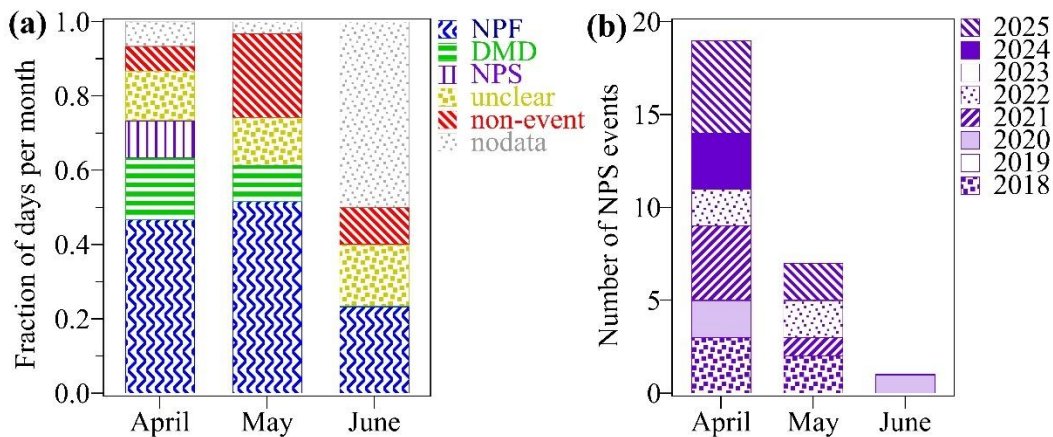
### ***2.2.8 Ancillary observations***

Collocated (within a 50 m radius) trace gas measurements, including NO<sub>x</sub>, CO, and O<sub>3</sub>, as well as meteorological parameters, including air temperature, relative humidity, solar radiation, wind speed, and wind direction, were obtained from the national air quality monitoring network operated by the DLI of Cyprus. O<sub>3</sub> was measured with UV photometric monitors (Ecotech 9810B). CO was measured with non-dispersive IR (NDIR) spectroscopy monitors (Ecotech 9830B). NO<sub>x</sub> was measured with the chemiluminescence technique. Solar radiation was measured with a Kipp & Zonen CMP22 solar irradiance sensor. Further details of DLI instrumentation are available in Vrekoussis et al. (2022). In addition, aerosol optical depth (AOD) was obtained from the AERONET sun photometers operated at the CAO-AMX site. All data are reported in Coordinated Universal Time (UTC). Local time in Cyprus is UTC+3 during daylight saving time (Eastern European Summer Time, late March to late October). Only quality-assured and quality-controlled datasets, based on standard procedures and instrument-specific data quality flags, were used in this study. Air-mass backward trajectories were computed using NOAA’s Air Resources Laboratory (ARL), Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) PC-version model, version 4.4 (Draxler and Rolph, 2010; Stein et al., 2015), incorporating meteorological data from the Global Data Assimilation System (GDAS) with a spatial resolution of 0.25° × 0.25° and a temporal resolution of 1 hour (Draxler and Rolph, 2010). Two-day hourly backward air-mass trajectories, initialized at 500 m above the ground level, were used in this analysis. We further used ERA5, a fifth-generation reanalysis produced by the European Centre for Medium-Range Weather Forecasts (ECMWF), 0.25° gridded data of vertical velocity and specific humidity at hourly resolution on pressure levels from 1000 hPa to 550 hPa (Hersbach et al., 2023).

## **3. Results and discussion**

### ***3.1 Identification of event types***

Event types were classified by visual inspection of particle size distribution contour plots (Dal Maso et al., 2005). A day was identified as an NPF event day when a pronounced increase in particle number concentrations was observed in the nucleation size range (<25 nm), followed by continuous growth of these particles to larger diameters. Such events typically extend over spatial scales of hundreds of kilometers while they persist for 1–2 days, and are therefore characterised as regional NPF events (Kulmala et al., 2004). NPF events are often accompanied by a decrease in particle mode diameter from the Aitken size range during the afternoon hours, a phenomenon referred to as DMD (or shrinkage event) in the literature. As described earlier, we also observed events characterised by a rapid decrease in particle mode diameter within the sub-20 nm size range that occurred in the absence of the preceding particle formation and growth typically associated with DMD events, referred to as NPS. Such behavior has previously been reported at our site (Baalbaki et al., 2021) and in Pune, India (Kamra et al., 2022), but the underlying processes were not examined, and similar events have not been reported elsewhere. Notably, the particle size distribution contour plot during an NPS event displays a mirror image of the conventional “banana-shaped” NPF pattern, appearing as a distinctive “reverse-NPF” shape. A day without the onset of a new particle mode in the nucleation size range is classified as a non-event day. A day that could not be classified into any of the above categories was classified as an unclear day. A day without a valid full-day measurement, due to instrument maintenance, troubleshooting, or occasional power outages, was classified as a “nodata” day. Figure S1 shows the temporal evolution of median particle size distributions from nano Condensation Nucleus Counter (nCNC), Neutral Air Ion Spectrometer (NAIS; negative polarity ions and particles), and scanning mobility particle sizer (SMPS), while Fig. S2 shows typical examples of NPF, DMD, NPS, unclear, and non-event days.



**Figure 2:** (a) Monthly fraction of days classified into different event types during the SPICY campaign (1 April – 15 June 2024). (b) Total number of NPS event days identified from 2018 to 2025. There are no measurements during 2019 and 2023.

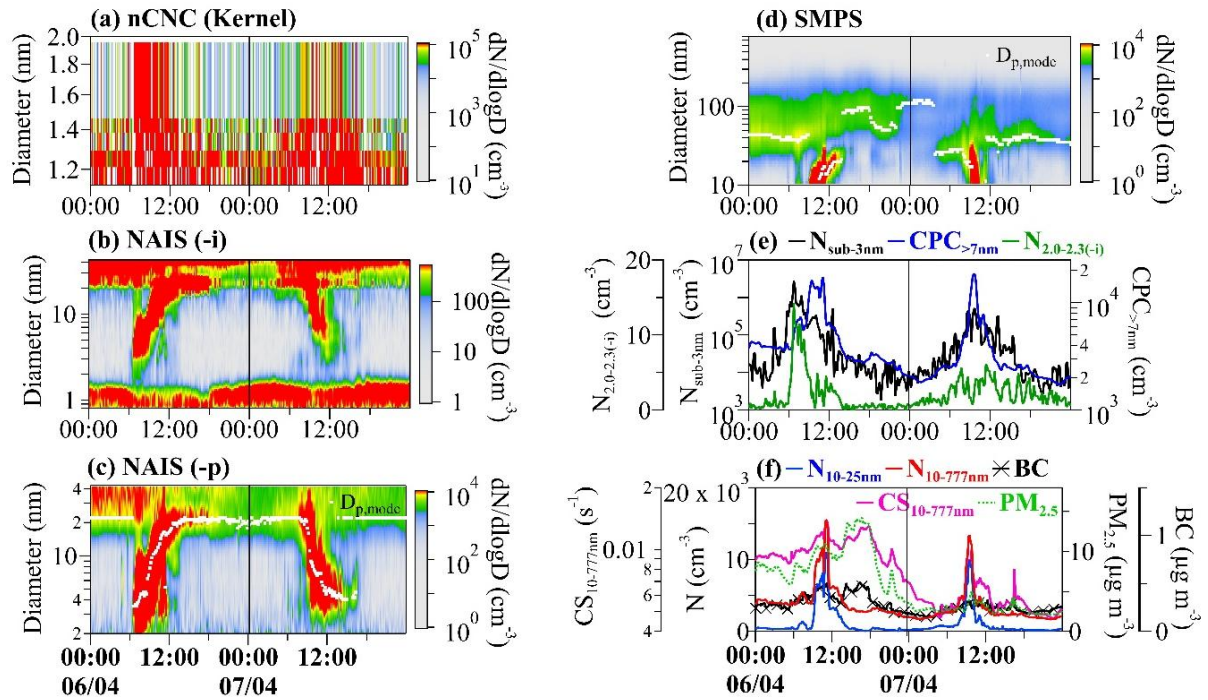
Figure 2a shows the monthly fraction of days classified into different event types during the SPICY campaign. NPF events accounted for the largest fraction of days, occurring on 44 of the 76 campaign days (57.9%). Of these, 37 days were classified as conventional NPF events (48.7%), while 8 days were identified as DMD events (10.5%). NPS events were observed on 3 days (3.9%). non-events and unclear days accounted for 12 (15.8%) and 13 days (17.1%), respectively. There were no valid measurements on 3 days (3.9%) during the campaign. Here, we present a two-day consecutive case study consisting of an NPF event on 6 April followed by an NPS event on

255 7 April. This consecutive case was selected to minimize variability associated with regional meteorological conditions and to enable a detailed analysis of the physicochemical mechanisms underlying the NPS phenomenon. In addition, two further consecutive NPF-NPS cases (11-12 April and 20-21 April) are presented in the Supplementary Information (Figs. S3-S7), and all three cases are summarised below. Long-term measurements of particle size distributions at the CAO-AMX site indicate that NPS is a commonly observed phenomenon (Fig. 260 2b), highlighting the importance of the comprehensive SPICY observations for elucidating its underlying mechanisms.

### ***3.2 Elucidating a nanoparticle shrinkage event***

#### ***3.2.1 Particle size distributions and aerosol properties during consecutive NPF and NPS events***

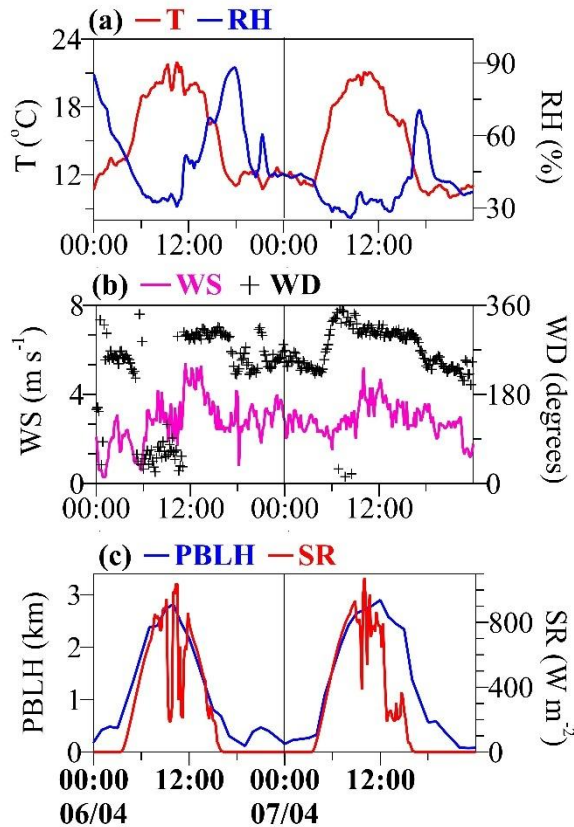
265 The median diurnal variation of particle size distributions and aerosol properties during two consecutive events, an NPF event on 6 April followed by an NPS event on 7 April, are shown in Fig. 3. A sharp increase in sub-3 nm particle number concentrations (nCNC; Fig. 3a), followed by a typical “banana-shaped” aerosol growth pattern in the particle size distribution (Fig. 3b-d), was observed during the NPF event. In contrast, the NPS event exhibited a rapid decrease in nanoparticle mode diameter (<20 nm) without any preceding evidence of particle formation or 270 growth. During the NPF event, an approximately 2-hour time lag between the peak number concentrations of sub-3nm particles (nCNC) and >7 nm particles (CPC) clearly indicate particle growth, whereas the nearly simultaneous peaks observed during the NPS event suggest the absence of particle growth (Fig. 3e). While the concentrations of ions with diameters 2.0 - 2.3 nm, indicative of local intermediate ion (2 - 7 nm) formation (Tuovinen et al., 2024), increased sharply during the NPF event, they remained negligible during the NPS event 275 (Fig. 3e). This indicates the absence of local NPF processes during the NPS event. The number concentrations of nucleation mode particles ( $N_{10-25\text{nm}}$ ) and total particles ( $N_{10-777\text{nm}}$ ) were comparable on 6 and 7 April. In contrast, both the total condensation sink ( $CS_{10-777\text{nm}}$ ) and particulate matter with an aerodynamic diameter less than 2.5  $\mu\text{m}$  ( $\text{PM}_{2.5}$ ) were approximately twofold lower during the NPS event than during the NPF event (Fig. 3f). While a lower CS generally favors the growth of small clusters or newly formed particles by reducing vapour losses to pre-existing particles (Kulmala et al., 2001), it also indicates cleaner atmospheric conditions with a limited 280 availability of condensing vapours. Consistently, black carbon (BC) mass concentrations were slightly lower during the NPS event, suggesting a minimal influence from local anthropogenic sources such as fossil fuel combustion or biomass burning. This further suggests the absence of anthropogenic sources of nanoparticles upwind of the site. Two other consecutive NPF-NPS cases (10-11 April and 20-21 April) showed a similar 285 temporal evolution of particle size distributions and aerosol properties (Figs. S3 and S4), except for higher pre-existing particle concentrations during the NPS event in the 10-11 April case.



290 **Figure 3:** Median diurnal variation of aerosol properties during the consecutive NPF (6 April) and NPS (7 April)  
 events. Particle size distributions measured by (a) nCNC, (b-c) NAIS negative polarity ions and particles, and (d)  
 SMPS. Concentrations of (e) negative polarity ions in the 2.0-2.3 nm size range measured by NAIS ( $N_{2.0-2.3(-i)}$ ),  
 sub-3nm particles measured by nCNC ( $N_{\text{sub-3nm}}$ ), and >7nm particles measured by CPC ( $N_{>7\text{nm}}$ ); (f) nucleation  
 295 mode ( $N_{10-25\text{nm}}$ ) and total particle ( $N_{10-777\text{nm}}$ ) number concentrations from SMPS, total condensation sink ( $CS_{10-777\text{nm}}$ ), and particulate matter of aerodynamic diameter less than 2.5  $\mu\text{m}$  ( $PM_{2.5}$ ) and black carbon (BC) mass  
 concentrations.

### 3.2.2 Influence of meteorological conditions and air-mass transport

300 Meteorological conditions were largely comparable during the NPF event on 6 April and the NPS event on 7 April  
 (Fig. 4). The diurnal temperature variation was similar, with mean temperatures of 20.1  $^{\circ}\text{C}$  and 19.9  $^{\circ}\text{C}$  (averaged  
 over 06-12 UTC) during the NPF and NPS events, respectively. Relative humidity also showed little difference,  
 with mean values of 30.4% during the NPF event and 30.7% during the NPS event. Wind speeds were similar for  
 both events, although wind direction differed, being northeasterly on 6 April and northwesterly on 7 April during  
 the same period (Fig. 4b). Boundary layer height and solar radiation (Fig. 4c), which are key factors influencing  
 305 new particle formation and growth (Kulmala et al., 2004; Deot et al., 2025), likewise exhibited no substantial  
 differences between the two events. Overall, local meteorological conditions were similar during the NPF and  
 NPS events, with wind direction being the notable difference. A comparable temporal evolution of meteorological  
 parameters was also observed for the two additional consecutive NPF-NPS cases (Fig. S5), suggesting that the  
 contrasting particle behavior was unlikely to be primarily driven by local meteorological variability but rather by  
 310 variations in air-mass history (discussed next).

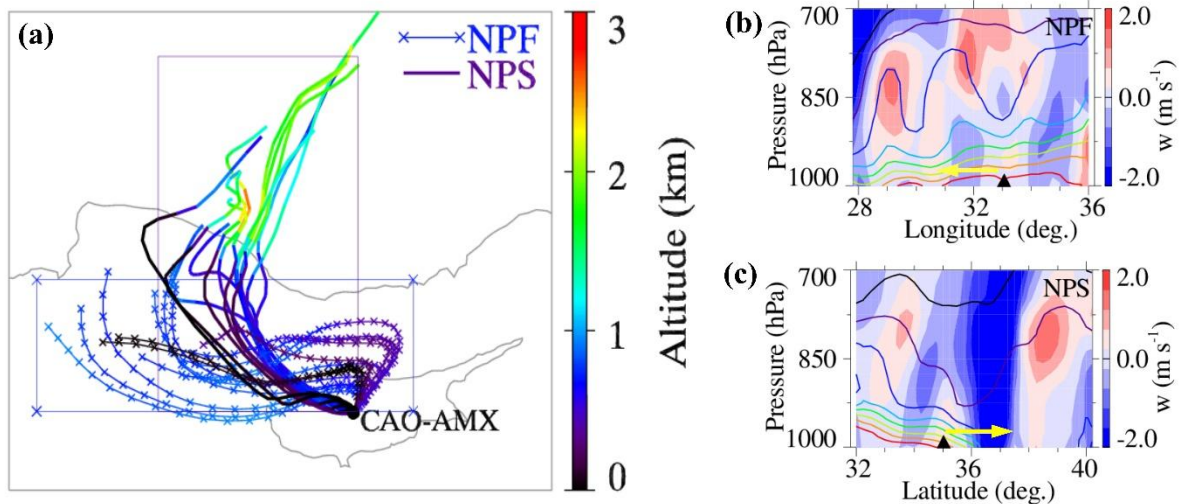


**Figure 4:** Median diurnal variation of meteorological parameters during the consecutive NPF-NPS events (6-7  
 315 April). (a) air temperature (T) and relative humidity (RH), (b) wind speed (WS) and wind direction (WD), and (c)  
 planetary boundary layer height (PBLH) and solar radiation (SR).

Previous studies have shown that variations in air-mass origin, transport pathways, atmospheric dilution, vertical  
 mixing, and the mixing of different air-masses can strongly influence the occurrence and evolution of NPF events  
 320 (Kulmala et al., 2004; Kanawade et al., 2012; Nilsson et al., 2001; Hussein et al., 2009; Tunved et al., 2004). Air-  
 mass trajectory analysis reveals distinct transport pathways and air-mass characteristics between the two cases,  
 particularly on 6 and 7 April (Fig. 5a). During the NPF event, air masses traveled slowly near the ocean surface  
 and largely remained within the PBL. In contrast, during the NPS event, the air masses traveled fast and descended  
 from the free troposphere to near-surface levels during the passage over the Western Taurus mountain range (Fig.  
 325 5a). A similar contrast in air-mass history between NPF and NPS events was observed for the 10-11 April case  
 (Fig. S6a). While the air masses traveled within the PBL during both the NPF and NPS events for the 20-21 April  
 case (Fig. S7a), air masses traveled fast near the ocean surface on 21 April (NPS event).

To further assess the dynamic state of the lower atmosphere, vertical velocity and specific humidity were  
 330 examined (Fig. 5b-c). Stronger low-level subsidence, accompanied by lower specific humidity (i.e., drier air), was  
 evident in the upwind region of the measurement site on NPS event days (Fig. 5c), indicating the intrusion of drier  
 air masses from higher altitudes. Such enhanced vertical mixing and entrainment of cleaner, drier free tropospheric  
 air are indicative of atmospheric dilution and are consistent with the observed lower near-surface aerosol mass  
 (Fig. 6a-b) and reduced columnar aerosol loading (Fig. 6c), as well as lower  $\text{NO}_x$  (Fig. 6d) and carbon monoxide

335 (CO; Fig. 6e). Fast-moving air-masses accompanied by strong winds can enhance turbulent mixing, leading to dilution of particle concentrations and changes in particle size distributions (Shi et al., 1999), which is evident during NPS events (Fig 5a and Figs. S6-S7). Furthermore, the observed nucleation mode particles may originate from NPF occurring upwind of the measurement site and subsequently advected to the site, or from overlapping nucleation modes arising from distinct nearby anthropogenic sources (Hakala et al., 2023; Kivekäs et al., 2016).  
 340 As discussed earlier, anthropogenic sources upwind of the measurement site are minimal (Fig. 3f); thus, the advection of nucleation-mode particles directly from primary emissions is unlikely to explain the observed nucleation mode particles during the NPS event.

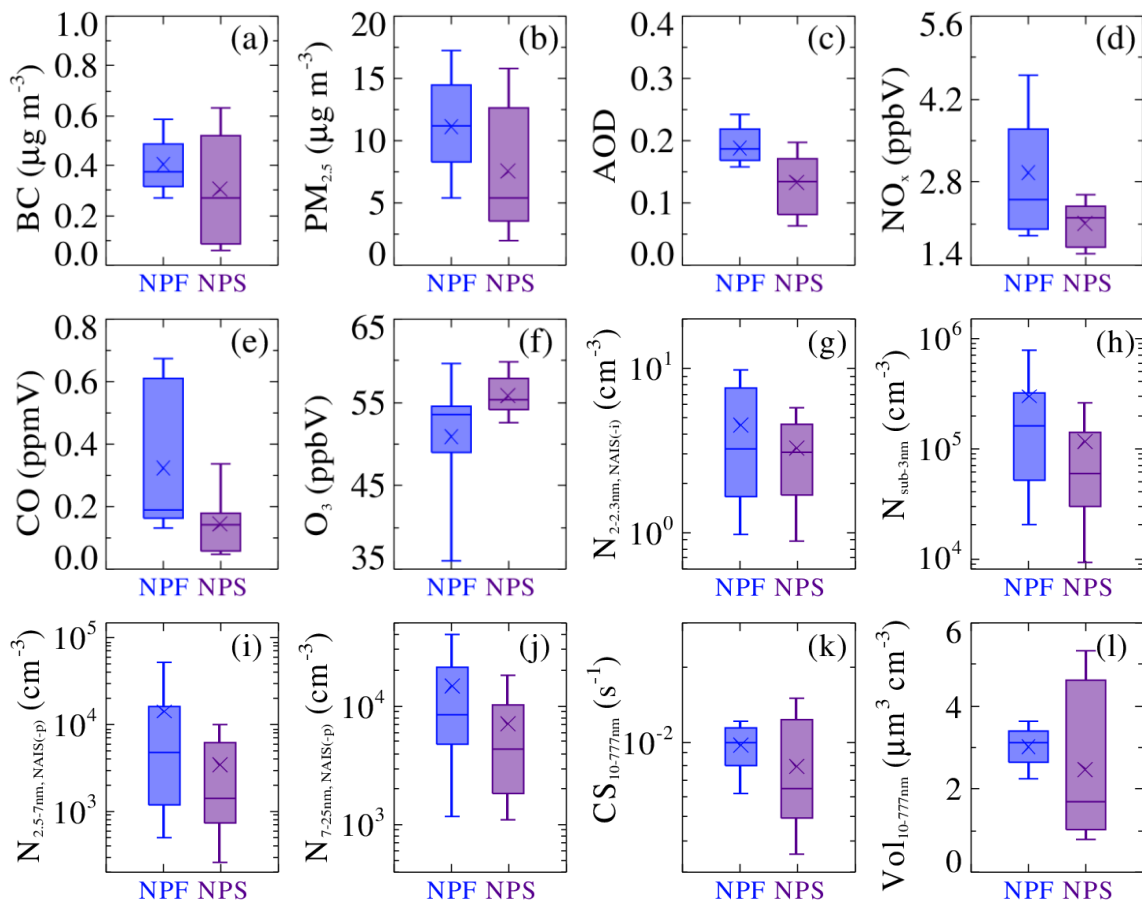


**Figure 5:** 48-hour air-mass backward trajectories (00-12 UTC) as a function of altitude, initialized at 500 meters above ground level at the AMX site during the NPF event (6 April, line connected by cross symbol) and the NPS event (7 April, solid lines). The region shown by the rectangular box is used to create longitude (latitude)-altitude cross-sections of averaged (b-c) vertical velocity ( $w$ , filled contours) along air-mass trajectories over 0-9 UTC for NPF and NPS event days observed on 6 April and 7 April, respectively. Contour lines represent specific humidity of 2, 3, 4, 5, 6, 7, 8, and 10  $\text{g kg}^{-1}$ , shown in black, violet, blue, cyan, green, yellow, orange, and red, respectively.  
 350 The negative value of  $w$  indicates subsidence, while the positive values indicate the updraft. The yellow-colored arrows indicate the upwind region of the measurement site.

The origin of the observed nanoparticles during NPS event days can be constrained into three physically consistent scenarios: the particles were either transported from nearby anthropogenic sources or formed upwind of the measurement site and subsequently advected to the site or formed locally but failed to grow. The low BC mass concentrations (Figs. 3f, 6a), together with reduced  $\text{NO}_x$  (Fig. 6d) and CO (Fig. 6e) levels, suggest that contributions from overlapping nucleation modes arising from nearby anthropogenic sources (Hakala et al., 2023; Kivekäs et al., 2016) are unlikely to explain the observed nucleation mode particles during the NPS event. Instead, these particles are more likely to be formed upwind of the site and subsequently transported to the observation  
 360 location. It is therefore critical to constrain the spatial extent of over which particle formation occurs. Two approaches are commonly used to estimate this extent. The first, and more resource-intensive, approach relies on simultaneous measurements of particle size distributions at multiple stationary sites along the air-mass transport pathway. The second approach is based on hourly calculated backward air-mass trajectories. As no simultaneous

365 measurements were available at the upwind locations of the measurement site, the spatial extent of the observed NPF and NPS events was estimated based on backward air-mass trajectory analysis, following the methodology of Hussein et al. (2009). This analysis reveals similar spatial scales for both event types (NPF and NPS; approximately 50-200 km, Fig. S8), with NPS events occurring along similar air-mass path. This suggests that particle formation occurring upwind of the measurement site on NPS event days is plausible. In our recent study, using simultaneous measurements from a low-altitude site (CAO-AMX) and a high-altitude site (Troodos, CAO-TRO), we showed that approximately half of the observed NPF events at both sites occurred concurrently along the same air-mass pathway, under the influence of an evolving planetary boundary layer height (Deot et al., 2025). The lower number concentrations of negative ions in 2.0-2.3 nm size range (Fig. 6g), sub-3nm neutral particles (Fig. 6h), and negative particles in the 2.5-7 nm (Fig 6i) and 7-25 nm (Fig 6j) size range on NPS event days can be attributed to reduced levels of condensing vapours (discussed in Section 3.2.3), together with a lower condensation sink (Fig. 6k) and a lower particle volume concentration (Fig. 6l).

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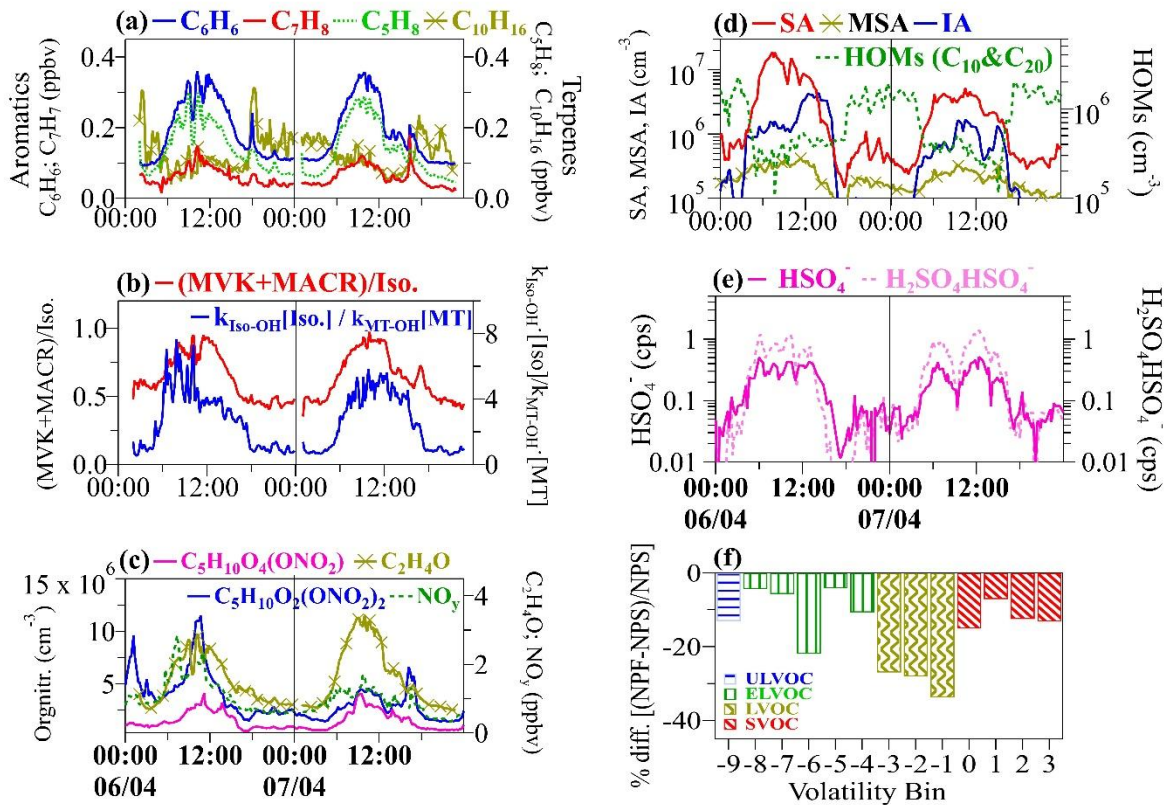
380 **Figure 6:** Key differences in aerosols, gases, and nanoparticles averaged over the observed NPF (6, 10, and 20 April) and NPS (7, 11, and 21 April) event days. Box-whisker plots show (a) black carbon (BC) mass concentrations, (b) particulate matter with an aerodynamic diameter less than 2.5  $\mu\text{m}$  ( $\text{PM}_{2.5}$ ), (c) aerosol optical depth (AOD), (d) oxides of nitrogen ( $\text{NO}_x$ ), (e) carbon monoxide (CO), (f) ozone ( $\text{O}_3$ ), (g) negative ions in the 2.0–2.3 nm size range, (h) sub-3nm particles, (i) negative particles in the 2.5–7.0 nm size range, (j) negative particles in the 7.0–25 nm size range, (k) total condensation sink ( $\text{CS}_{10-777\text{nm}}$ ), and (l) total particle volume concentration. Data correspond to the 06–12 UTC period. The cross symbol indicates the mean, the horizontal

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line indicates the median, the bottom and top of the box indicate the 25<sup>th</sup> and 75<sup>th</sup> percentiles, and the bottom and top of the whisker indicate the 10<sup>th</sup> and 90<sup>th</sup> percentiles.

### 390 3.2.3 Role of precursor gases and condensing vapours

395 Volatile organic compound (VOC) mixing ratios and their early oxidation products were examined to identify the dominant chemical regimes associated with NPF and NPS events. VOCs at the CAO-AMX site are controlled by local temperature and continental transport, with long-term observations indicating the onset of local biogenic emissions in mid-March each year. Mixing ratios of benzene and toluene (anthropogenic tracers), as well as biogenic tracers such as isoprene and monoterpenes (MTs), were comparable between the two days (Fig. 7a). Methyl vinyl ketone (MVK) and methacrolein (MACR), first-generation OH-initiated oxidation products of isoprene, showed similar behavior on both days; accordingly, the (MVK+MACR)/isoprene ratio, a proxy for the extent of isoprene oxidation, was also comparable (Fig. 7b). Plant chamber studies (Kiendler-Scharr et al., 2009) and field observations (Kanawade et al., 2011) have shown that isoprene can suppress biogenic NPF at ground level, thereby dampening aerosol cooling effects (Lee et al., 2016), with the extent of suppression depending on the relative contributions of isoprene- and monoterpene-derived carbon. To compare the relative OH-initiated oxidation rates of isoprene and MTs, we calculated the ratio  $(k_{\text{Iso-OH}} \times [\text{Isoprene}]) / (k_{\text{MT-OH}} \times [\text{MT}])$ , where  $k_{\text{Iso-OH}} = 1.01 \times 10^{-10} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$  and  $k_{\alpha\text{-pinene-OH}} = 5.36 \times 10^{-11} \text{ cm}^3 \text{ molecules}^{-1} \text{ s}^{-1}$  represent the reaction rate coefficients of isoprene and alpha-pinene with OH radical, respectively (Finlayson-Pitts and Pitts, 2000). This ratio was slightly higher during the NPF event (Fig. 7b), indicating an isoprene-dominated OH-reactivity regime that favors the formation of more volatile, lower oxygen-to-carbon (O/C) oxidation products, which are less favorable for biogenic NPF. In contrast, the lower ratio observed during the NPS event rules out isoprene-driven suppression of biogenic NPF under those conditions; nevertheless, NPF did not occur at the site on 7 April. Curtius et al. (2024) showed that organonitrates formed from OH-initiated oxidation of isoprene in the presence of nitrogen oxides drive NPF in the upper troposphere over the Amazon. Consistently, concentrations of isoprene-derived hydroxy hydroperoxy nitrate ( $\text{C}_5\text{H}_{10}\text{O}_4(\text{ONO}_2)$ ) measured by MION-APi-TOF were comparable on both days, indicating that the initial stages of isoprene oxidation proceeded similarly during the two events (Fig. 7c). In contrast, peak concentrations of isoprene-derived dinitrates ( $\text{C}_5\text{H}_{10}\text{O}_2(\text{ONO}_2)_2$ ) were slightly higher during the NPF event than during the NPS event, indicating a stronger influence of high- $\text{NO}_x$  driven chemistry during the NPF event. This interpretation is supported by the reactive nitrogen compounds ( $\text{NO}_y$ ) concentrations, which were also slightly higher during the NPF event than during the NPS event (Fig. 7c). Further, acetaldehyde ( $\text{C}_2\text{H}_4\text{O}$ ), a marker of atmospheric oxidation, also exhibited similar mixing ratios on both days, indicating comparable overall oxidation levels (Fig. 7c).



**Figure 7:** Median diurnal variation of aerosol precursors and condensing vapours during the consecutive NPF-NPS events observed during 6-7 April. (a) mixing ratios of benzene ( $C_6H_6$ ), toluene ( $C_7H_8$ ), isoprene ( $C_5H_8$ ), and monoterpenes ( $C_{10}H_{16}$ ); (b) the ratio of the sum of methyl vinyl ketone (MVK) and maethacrolein (MACR) to the isoprene [(MVK+MACR)/Iso.] and the relative OH reaction rates of isoprene and MTs ( $k_{Iso-OH}[Iso]/k_{MT-OH}[MT]$ ); (c) organonitrates [ $C_5H_{10}O_4(ONO_2)$  and  $C_5H_{10}O_2(ONO_2)_2$ ], acetaldehyde ( $C_2H_4O$ ) and reactive nitrogen compounds ( $NO_y$ ); (d) concentrations of sulfuric acid (SA), methyl sulfonic acid (MSA), iodic acid (IA), and highly oxygenated organic molecules (HOMs,  $C_{10}$  &  $C_{20}$ ); (e) ambient ion mode signals of bisulfate monomer ( $HSO_4^-$ ) and dimer ( $H_2SO_4HSO_4^-$ ) in counts per second (cps); (f) percentage difference in organic vapour concentrations (molecules  $cm^{-3}$ ) between the NPF and NPS events as a function of volatility bin, expressed as  $\log_{10}$  of the effective saturation concentration,  $\log_{10}(C^*)$ . Negative percentage differences indicate higher organic vapour concentrations during the NPS event. ULVOC, ELVOC, LVOC, and SVOC denote ultra-low volatile organic carbon, extremely low volatile organic carbon, low volatile organic carbon, and semi-volatile organic carbon, respectively.

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Next, we examined neutral aerosol precursors, including sulfuric acid (SA), methanesulfonic acid (MSA), iodic acid (IA), and HOMs, measured using the nitrate mode of the MION-API-TOF. SA concentrations were approximately five-fold higher during the NPF event than during the NPS event (Fig. 7d). Nevertheless, SA concentrations during the NPS event ( $\sim 4 \times 10^6$  molecules  $cm^{-3}$ ) were sufficient to trigger NPF under atmospherically relevant ammonia mixing ratios of 100 parts per trillion by volume (pptv) and dimethylamine mixing ratios of 40 pptv (Kirkby et al., 2011; Kürten et al., 2018). Concurrently, HOMs concentrations increased during the NPF event, but decreased during the NPS event (Fig. 7d), whereas MSA and IA concentrations were

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comparable on both days (Fig. 7d). Negative ion measurements from APi-TOF further showed that concentrations of bisulfate monomers ( $\text{HSO}_4^-$ ) and dimers ( $\text{H}_2\text{SO}_4\text{HSO}_4^-$ ) were also similar on both days (Fig. 7e), indicating comparable levels of ionised SA clusters in the ambient air. No indication of higher multimers or ammonia-containing clusters was detected.

Chamber experiments have demonstrated that initial nanoparticle growth is dominated by extremely low-volatility organics (ELVOC; approximately  $C^* < 10^{-4.5} \mu\text{g m}^{-3}$ , volatility bin  $< -4$ ), whereas growth at larger sizes is increasingly supported by more abundant, slightly higher-volatility vapours in the low-volatility organic compounds (LVOC) range ( $C^* \approx 10^{-4.5} - 10^{-0.5}$ ;  $-4 > \text{volatility bin} < 0$ ) as the Kelvin barrier falls with increasing diameter (Tröstl et al., 2016). We, therefore, examined differences in organic vapour concentrations relevant to early cluster formation and subsequent growth between the two event types (Fig. 7f). Organic vapour concentrations were higher during the NPS event than during the NPF event across all volatility bins (Fig. 7f), indicating that particle growth was not limited by the availability of condensable vapours. Instead, these conditions point to an environment less favorable for stable initial cluster formation and subsequent growth, and more favorable to net evaporation of condensed material. Fast-moving air-masses enhance turbulent mixing, leading to dilution of particle concentrations, precursor gases, and changes in particle size distributions (Shi et al., 1999). This dilution lowers organic aerosol concentrations and reduces the absorptive capacity of the particle phase. As a result, gas-particle equilibrium shifts towards the vapour phase, enhancing evaporation of semi-volatile species from particle surfaces to re-establish equilibrium (Pankow, 1994; Zhang and Wexler, 2004). This repartitioning can cause aerosol mass to decrease more rapidly than expected from dilution alone, reflecting the dynamic response of SVOCs to changing ambient conditions (Donahue et al., 2014). Under ambient conditions, SVOCs partition between the gas and particle phases, whereas LVOCs and ELVOCs exist primarily, or entirely, in the particle phase (Barsanti et al., 2017). Consistent with this framework, the observed reduction in HOM concentrations (Fig. 7d, Figs. S9-S10) and the absence of particle growth suggest that dilution associated with the entrainment of cleaner, drier free tropospheric air into the boundary layer (Fig. 5, Figs. S6-S7) played a key role during NPS events. The similar chemical environment (Fig. 7; Figs. S9-S10) and local meteorological conditions (Fig. 4; Fig. S5), together associated with the contrasting particle evolution (Fig. 3; Figs. S3-S4), indicate that the observed differences in particle behavior were not primarily driven by low condensing vapours but are instead associated with atmospheric dilution.

The higher concentrations of LVOCs and SVOCs observed during the NPS event (Fig. 7f) would, in principle, be expected to condense onto existing particles and promote growth; however, condensation onto the smallest particles is likely suppressed by the strong curvature (Kelvin) effect. This inhibition limits nanoparticle growth despite the availability of condensable material, preventing particles from growing to larger sizes. Consistent with this interpretation, the low total particle volume concentrations observed during NPS event days (Fig. 6l), together with elevated LVOC and SVOC concentrations (Fig. 7f), point to evaporation of condensed material primarily driven by atmospheric dilution.

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#### 4. Conclusions

This study presents new evidence of nanoparticle shrinkage observed at the rural background site (CAO-AMX) in Cyprus. These events are characterised by a rapid reduction in particle diameters within the sub-20 nm size range, with the particle size distributions contour plot displaying a mirror image of the conventional “banana-shaped” NPF pattern. To our knowledge, this is the first time such a behavior is reported and analyzed, expanding the current phenomenology of atmospheric aerosol evolution.

Using a suite of instruments that enabled comprehensive gas- and particle-phase measurements, we conclude that NPS events are not primarily driven by low vapour concentrations but associated with atmospheric dilution, as indicated by air-mass trajectory analysis. SA concentrations during NPS events were sufficient to trigger NPF under atmospherically relevant boundary layer conditions, while key VOC mixing ratios and oxidation rates were comparable between NPF and NPS cases. Moreover, the limited local cluster formation, low black carbon and NO<sub>x</sub> concentrations, and a distinct air-mass source region rule out local primary emissions as the origin of the observed sub-20 nm particles. The similar spatial scales of these events suggest that the sub-20nm particles likely originated from particle formation occurring upwind of the measurement site. These events were further characterized by fast-moving air masses, which can enhance turbulent mixing, leading to dilution of particle concentrations and changes in particle size distributions. Such conditions reduce organic aerosol mass and shifts gas-particle equilibrium towards evaporation. Volatility-resolved analysis further indicated a dominance of organic compounds of low and moderate volatility. Although ULVOCs and ELVOCs concentrations were slightly higher during NPS events, particle formation and subsequent growth remained inhibited, supporting net evaporation and leading to rapid particle shrinkage.

Our findings highlight a previously unrecognised sink for nanoparticles occurring under conditions favorable for particle formation and growth. This can negatively impact the CCN budget, thereby damping the aerosol-induced radiative cooling effect.

#### **Code and data availability**

Time-synchronised 10-minute median 1D and 3D data variables are available at Zenodo (<https://doi.org/10.5281/zenodo.18685092>). Ceilometer data can be viewed at <https://e-profile.eu/>. ERA5 reanalysis data are publicly available from Copernicus Climate Change Service (C3S) Climate Data Store (CDS) at <https://cds.climate.copernicus.eu/datasets> (last access: January 2025). AERONET aerosol optical depth data are available publicly to download from <https://aeronet.gsfc.nasa.gov/> (last access: October 2025). Surface elevation data from the U.S. Geological Survey Earth Resources Observation and Science (EROS) Center is available publicly to download from <https://www.usgs.gov/centers/eros/science/usgs-eros-archive-digital-elevation-global-30-arc-second-elevation-gtopo30> (last access: 22 October 2024)

The IGOR Pro (V9) data analysis tool is available from the corresponding author upon reasonable request.

#### **Supplement link**

The link to the supplement will be included by Copernicus, if applicable.

### **Author contributions**

VPK, MP, FM, GB, EB, JS, and TJ designed the experiments. ND, AC, MvdB, AP, MP, SB, and CS carried them out. VPK, MC, AC, CL, AP, RB, MP, and AG analyzed the data. VPK prepared the manuscript with contributions  
525 from all co-authors.

### **Competing interests**

The author declares that they have no conflict of interest.

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### 540 **Acknowledgments**

This study was supported by the European Union ERC-2022-STGERC-BAE-Project: 101076311. This project has also received funding from the European Union's Horizon 2020 research and innovation programme (grant No. 856612) and the Cyprus Government (EMME-CARE). EB acknowledges support from the EVOCPOLIS project, funded by the Cyprus Research and Innovation Foundation (Grant No. Vision ERC-Parth 2/0524/0215).

545 The authors thank the University of Helsinki, INAR/Physics, and CiGas-UHEL for support and instrumentation. The authors also thank the technical support team: Nikoleta Lekaki, Moreno Parolin, Pierre-Yves Quehe, and Rafail Konatzii for maintaining field stations. The authors acknowledge valuable comments and suggestions from the reviewers.

### 550 **Funding Support**

This work received funding from the European Union ERC-2022-STGERC-BAE-Project: 101076311, the European Union's Horizon 2020 research and innovation programme (grant No. 856612) and the Cyprus Government (EMME-CARE) and the EVOCPOLIS project, funded by the Cyprus Research and Innovation Foundation (Grant No. Vision ERC-Parth 2/0524/0215).

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