# CIAO observatory main upgrade: building up building-up an

# 2 **ACTRIS compliant aerosol in-situ laboratory**

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#### 12 Abstract

13 This paper describes the aerosol in-situ laboratory at CIAO (CNR-IMAA Atmospheric Observatory) the CNR-IMAA 14 (Consiglio Nazionale delle Ricerche – Istituto di Metodologie per l'Analisi Ambientale) Atmospheric Observatory (CIAO) in 15 South Italy, outlining its configuration and detailing each instrument and sampling lines. The CIAO observatory has been 16 collecting observations of atmospheric components since 2000. Initially the activities revolved around aerosol lidar, later 17 radiosounding and cloud remote sensing observations were added over the years and made CIAO a leading atmospheric 18 observatory in the Mediterranean region. In 2018, a significant upgrade started for enhancing the observational capability by 19 adding aerosol in-situ instruments, with the objective to push new research boundaries for aerosol particles characterization 20 and multi-instrumental synergistic combined approaches. Here, we describe each technical implementation step for building 21 up an extensive aerosol in-situ laboratory compliant with ACTRIS (Aerosol Clouds and Trace gases Research InfraStructure) 22 standard operating procedures. Starting from scratch, the long path initiated in 2018, with the design of the laboratory in terms 23 of instruments, container organization layout, inlets and sampling lines optimizations, that required time and interactions with 24 experts in the field. Reporting here all the details about the final solutions implemented at CIAO, this paper will be, for new 25 aerosol in-situ laboratory, a practical guide for the implementation of the aerosol in-situ observational site.

#### 26 1 Introduction

27 The importance of a quantitative and qualitative assessment of atmospheric aerosol particle characteristics has been recognized 28 since many years; aerosole particles are responsible for direct and indirect effects on atmospheric processes, affecting climate 29 and human health, as well as precipitation cycle and air quality (e.g. Pöschl, 2005; Ren-Jian et al., 2012; IPCC, 2021). 30 Depending on their sources, aerosols particles appear in different sizes/shapes and their relatively short lifetime makes the 31 physical and chemical properties extremely variable both on temporal and spatial scales. Because of the inherent complexity 32 of aerosols particles, a single measurement technique providing all the relevant information is not available: thus, a multi-33 instrument approach is needed. The combination of different techniques and observational platforms can be crucial for a better 34 understanding of the presence and the characteristics of atmospheric aerosols particles, as well as their role in the large variety 35 of processes in which they are involved. The Aerosol Clouds and Trace Gases Research InfraStructure (ACTRIS, 36 www.actris.eu; Laj et al., 2024) is the European--Research Infrastructure (E-RI) aiming to integrate previous existing networks 37 for the characterization of aerosols particles, clouds and trace gases using and integrating in-situ and remote sensing 38 observations, and experimental platforms for the characterization of atmospheric components under controlled environments. 39 An overarching investigation of the atmosphere which accounts for all these three components is a winning strategy: For 40 instance, aerosols particles act as cloud condensation nuclei (CNN) affecting the cloud properties and lifetime; emitted gas 41 species may act as precursors to form new particles in the atmosphere, i.e., the secondary aerosol particle. Integrated 42 approaches of remote sensing and in-situ observations allows to take the most from the detailed and accurate characterization 43 in terms of morphology of particles, dimension and chemical composition: the remote sensing provide the vertical profile of 44 physical and optical properties information which are essential for investigating aerosol layers, long range transportation, 45 mixing processes and aerosol-cloud interactions; the latter is the only approach to provide the chemical composition and 46 reliable data at ground level, where aerosols particles affect ecosystems and humans.

47 In this scenario, the CNR-IMAA (Consiglio Nazionale delle Ricerche - Istituto di Metodologie per l'Analisi Ambientale) 48 Atmospheric Observatory (CIAO; Madonna et al., 2011), operating since 2000, has been recently upgraded with the aerosol 49 in-situ observational component, thus complementing the multi-year high-quality aerosol remote sensing data record. The 50 combination of the aerosol in-situ measurements with remote sensing observations is expected to strengthen fundamental 51 knowledge about aerosol particle impact on human health, ecosystems, and climate. This combination can be achieved either 52 by comparing or complementing the techniques: the results of the comparison will allow to reduce the uncertainty of aerosol 53 particles measurements in the atmosphere, with a subsequent improvement of model predictions on climate change, whereas 54 the complementarity results in the possibility of investigating the aerosol particles from the ground up to the stratosphere. The 55 new aerosol in-situ facility at CIAO, funded by the Italian Ministry of University and Research through the PER-ACTRIS-IT 56 project (https://www.imaa.cnr.it/en/projects/38-attivita/progetti/713-per-actris-it, last access: 6 December 2023), has received

- 57 initial acceptance as ACTRIS National Facility observational platform for the measurements of at least the obligatory ACTRIS 58 aerosol in-situ variables. The site has begun the next phase of the labelling process in 2024, a key element of ACTRIS's data 59 quality assurance system. This process ensures that instruments, data, and methodologies used across ACTRIS observational 60 platforms meet specific quality criteria. The labelling process involves a series of evaluations and certifications to verify 61 compliance with ACTRIS protocols (Deliverable 5.1; ACTRIS NF Labelling Plan). During the labelling process, the National 62 Facilities are annually invited by CAIS - ECAC (Center for Aerosol In-Situ - European - Center for Aerosol Calibration and 63 Characterization) to calibration workshops, where instruments are calibrated, and the quality of the data is thoroughly verified. The site will start the next phase of the labelling process in 2024. 64 65 In this paper we present a concise overview of the observatory, focusing on the characteristics of the recently established ACTRIS-compliant in-situ facility, with the main aim to benefit the aerosol community providing a comprehensive and 66 67 detailed description of technical solutions for the implementation of such component. A guidance for building-up an ACTRIS 68 aerosol in-situ station, it is potentially of interest also for extra European/outside of ACTRIS community: the ACTRIS in situ
- 69 standards are in some way following the WMO/GAW (World Meteorological Organization/Global Atmosphere Watch;
- 70 WMO/GAW, 2018) ones, therefore the interest in technical solutions for an ACTRIS compliant in-situ instrumentations stays
- 71 not only with the stations potentially involved in ACTRIS. Additionally, new EU air quality directive will include some more
- 72 advanced stations where black carbon and ultrafine measurements should be collected. Therefore, solutions adopted for
- 73 collecting such measurements with ACTRIS standard could be of interest for air quality management networks for guarantying
- 74 the quality of the collected data.
- After a short description of CIAO and typical atmospheric conditions in Sect. 2, Sect. 3 reports about the remote sensing instrumentation currently operating at CIAO. Section 4 represents the core of this paper, providing the in-depth description of the in-situ facility with the detailed configuration of each instrument and sampling lines. Finally, Sect. 5 illustrates three scientific topics to be studied at CIAO with the synergistic combined deployment of aerosol in-situ and remote sensing measurements.
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#### 81 **2 Description of the site**

Equipped with state-of-the-art systems for remote-sensing and in-situ measurements of aerosol\_particles, CIAO
 (<u>https://ciao.imaa.cnr.it, last access:</u> 4 December 2023) is currently a reference observatory for atmospheric research in

84 Europea reference station for short-live atmospheric constituents in Italy and the Mediterranean. The site is located on the

- 85 Southern Apennine in Italy (Tito Scalo, 40.60° N, 15.72° E, 760 m a.s.l.), in a plain surrounded by low mountains, less than
- 86 150 km away from the West, South and East coasts (Fig. 1).



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89 Figure 1: Location and image of CNR-IMAA Atmospheric Observatory (© Google Earth)



Figure 1: Location of the CNR-IMAA Atmospheric Observatory (© Google Earth); a) overview of the aerosol in-situ facility and
 b) overview of all CIAO infrastructures.

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Therefore, it operates in a typical mountainous weather strongly influenced by Mediterranean atmospheric circulation, resulting in generally dry, hot summers and cold winters. Indeed, dew point temperatures at the station between 20<u>1804</u> and 20<u>2117</u> after sunset exceeded 15-°C only during summer (Fig. 2). The prevailing wind direction occurring at the site is W-WSW-SW (Fig. 3).





100 Figure 2: Dew point temperature time series at CIAO in the timeframe 2018-2021 calculated from the RH and temperature values

101 measured at the ground level with the sensors of VAISALA RS41 radiosondes, typically launched twice a week between 30 and 120

102 minutes after sunset.



- 103
- Figure 2: Dew point temperature time series at CIAO in the timeframe 2004-2017 obtained from continuous measurements
   of the automatic weather station VAISALA MILOS520 with daily mean.
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107 Most of the surrounding land is classified as arable crops in non-irrigated areas, followed by broad-leaved woods and 108 coniferous forests, sclerophyllous or wooded/shrubby areas and natural grazing areas and grasslands 109 (http://rsdi.regione.basilicata.it, last access: 28 November 2023).

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#### 113 Figure 3: Wind rose diagram at CIAO in 2004-2017 obtained from continuous measurements of the automatic weather station

- 114 VAISALA MILOS520 with temporal resolution of 1 minute.
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116 CIAO's mission is to improve the knowledge of atmospheric processes and their role in meteorological phenomena, climate

and approaches adopted are the development of internationally recognized Standard Operating Procedures, the open data policy

change and air quality. Given the coverage and global relevance of the processes studied, fundamental aspects of the activities

- 119 and the full sharing of methodologies and know-how.
- CIAO provides free and open access to national and international users like researchers, <u>s</u>-mall and medium-sized enterprises (SMEs), students and citizens. At the present time, CIAO extends its outreach through the ATMO-ACCESS Trans-National Access program (https://www.atmo-access.eu/second-call-for-access/, last access: 2 December 2023). This program allows
- T22 Access program (<u>https://www.atmo-access.cu/second-can-tot-access/</u>last access. 2 December 2025). This program anows
- participants to engage in research on aerosols <u>particles</u> and their effects, to learn techniques and methods, and to contribute to
- 124 instruments, or to collaborate with the team.
- 125 The research activities of CIAO revolve around evolved within the long-term observations of aerosols particles, clouds, trace 126 gases and greenhouse gases within the European research infrastructures ACTRIS and ICOS (Integrated Carbon Observing 127 System), as well as around the participation of CIAO in reference observational programs and networks on a global scale, such 128 as GRUAN (GCOS (Global Climate Observing System) Reference Upper-Air Network) and GALION (GAW Aerosol Lidar 129 Observation Network). The observational strategy is organized to provide quality assured measurements data for satellite 130 validation and model evaluation and to fully exploit the synergy and integration of the active and passive sensors for the 131 improvement of the atmospheric characterization (e.g., Pappalardo et al., 2004;; Mona et al., 2009; Boselli et al., 2012; Ilić et 132 al., 2022). The complete list of CIAO publications is available at https://ciao.imaa.cnr.it/publications/.
- 133 For what concerns aerosol particles measurements, CIAO due to its geographical position as well as the low aerosol particles 134 background concentration is interesting for studying particles of natural origin such as desert dust and volcanic ash clouds. 135 The site is regularly affected by Saharan dust intrusions (e.g., Mona et al., 2006; Mona et al., 2014; Binietoglou et al., 2015; 136 Soupiona et al., 2020) has been reached by volcanic aerosol particles at the level of free troposphere during the eruptions of 137 Etna (e.g., Pappalardo et al., 2004a, Villani et al., 2006) and Eyjafjallajökull (Madonna et al., 2010; Mona et al., 2012; 138 Pappalardo et al., 2013) volcanos in 2002 and 2010, respectively, and stratospheric layers (e.g., Sawamura et al., 2012). In 139 recent years, the observatory has become actively involved in the study of smokes plumes originated by wildfires occurring 140 both at short-range, spreading with increased frequency in the surrounding forestry areas during the summer period (De Rosa 141 et al., 2022), and long-range transported plumes, such as the autumn 2020 California wildfires whose smokes transported in 142 the stratosphere reached the site within 13 days (Baars et al., 2019).
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#### 144 **3 Remote sensing measurements**

145 Remote sensing measurements have been the backbone of the research activity at CIAO since its beginning in the early 2000s, 146 with the scientific goal of providing long-term measurements for the climatology of aerosol and cloud properties. The main 147 research lines currently include; a) development of advanced lidar systems for the study of aerosols and aerosol cloud 148 interactions; b) design and implementation of new products (such as aerosol typing, aviation specific products, and 149 atmospheric boundary layer height); c) development and implementation of open data and FAIR data management policies 150 within the ARES node of the ACTRIS Data Center and RIs in the environmental field; d) development and implementation of 151 access policies to European RIs; e) deep/machine learning and signal processing applied to Earth observation; f) studies 152 integrated with transport models, satellite data and climate models; g) harmonization of the time series of measurements of 153 atmospheric variables; h) measurement campaigns for validation and integration with satellite data; i) networking at European 154 and global level. 155 Besides the compliance to the ACTRIS guidelines, all the remote-sensing measurements performed at CIAO are designed to

be in line with the main ground-based observation networks (i.e., EARLINET (European Aerosol Research Lidar Network to Establish an Aerosol Climatology), CloudNet, AERONET (AErosol RObotic NETwork), GRUAN, GALION) and the major international standards provided by the WMO/GAW 2016, aiming at establishing a long-term, harmonised and statistically significant database of measurements of atmospheric parameters and constituents for climatological studies (Matthias et al.,

160 2004).

161 The active remote-sensing instruments operative at CIAO include multi-wavelength Raman and polarization lidars, 162 ceilometers, Doppler lidars and polarimetric Doppler radars, and the passive ones include microwave radiometers, 163 photometers, and a high-resolution Fourier-Transform Infrared (FTIR) spectrometer.

With respect to the status of CIAO reported in previous papers (e.g., Madonna et al., 2011), some instruments are still operating,
 some have been replaced by more recent and advanced ones, and new instruments for increasing the observational capabilities
 have been added.

#### 167 The complete list of the remote sensing suite is reported in Table 1.

Aerosol Remote Sensing	Cloud Remote Sensing	Trace gases
		Remote Sensing
Fixed multi-wavelength Raman lidar	Ka-band Doppler radar Metek MIRA-35	
Mobile multi wavelength Raman lidar	Compact Ka band Doppler radar MIRA 35C	FTIR Bruker
Lidar and optical laboratories	W-band Doppler radar RPG-FMCW-94	<del>125HK</del>
	K-band Doppler radar Metek MRR-PRO	

MUSA Transportable Fixed multi-	Microwave radiometer RPG HATPRO G5	
wavelength Raman lidar		
Scanning UV Raman lidar	Ceilometer Vaisala CL51	
Automatic sun/sky/lunar photometer	Ceilometer Vaisala CL31	
Cimel 318T	Ceilometer Lufft CHM15k	
	2 Doppler lidars Halo Photonics Stream LineXR	

169 Table 1: List of the CIAO Remote Sensing instruments

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171 For the aerosol remote sensing, two new highly-advanced lidar systems have been recently installed at CIAO, one fixed and 172 one mobile. The first one will be an ACTRIS Observational Platform and the second one an Exploratory Platform available 173 even in combination with cloud remote sensing equipment. Both the lidars are capable of carrying out continuous 174 measurements going well beyond the required ACTRIS/EARLINET standards. They are able to provide measurements of 175 vertical profiles of several aerosol optical properties: backscatter coefficient and particle depolarization at 1064, 532 and 355 176 nm, and extinction coefficient at 532 and 355 nm, with the observational range starting from 200 m up to at least 20 km of 177 altitude. The fixed lidar is able to reach a measurement altitude range higher than 20km, being equipped with two telescopes, 178 and to provide vertical profiles of water vapor mixing ratio, this latter useful for investigating the impact of water vapour on 179 aerosol properties. On the other hand, the mobile system is more compact and transportable, and is used for field campaigns. 180 Both systems are part of the Centre for Aerosol Remote Sensing (CARS), one of the ACTRIS central facilities, that has the 181 mission to offer operation support to ACTRIS National Facilities operating aerosol remote sensing instrumentation. The two 182 systems are reference lidars for ACTRIS and offer services to test the performances of other lidar systems also through on-site 183 direct intercomparison campaigns using the mobile lidar.

184 Close by to the aerosol multiwavelength depolarization Raman, a triple mode photometer is operational within AERONET 185 (AErosol RObotic NETwork) and ACTRIS providing columnar aerosol optical depth measurements and columnar size 186 distribution information not only in daytime, but also in night-time under certain illumination conditions. CIAO is also 187 equipped with a lidar laboratory and an optical laboratory, which. The lidar laboratory is a facility that allows to implement 188 and test several and customized lidar configurations and (fluorescence, HSRL, multiwavelength elastic/Raman/depolarization 189 and water vapour mixing ratio and liquid water, and rotational Raman for temperature) in a modular way. The optical laboratory 190 allows to test and characterize optical components and laser sources typically used in high power lidar systems. Both 191 laboratories are also part of CARS and are open to users who want to benefit from the offered services.

- Besides the aerosol remote sensing instruments, cloud remote sensing equipment has also been updated and expanded with additional complementary instruments, setting up both an Observational Platform and a mobile Exploratory Platform
- 194 compliant with ACTRIS/CloudNet requirements. Among the complementary instruments, Doppler lidars are capable of

- 195 measuring the profiles of horizontal and vertical wind and related fluid dynamic parameters through the troposphere. Finally,
- 196 and-a high resolution FTIR spectrometer has been added for performing remote sensing measurements of trace gases to 197 complement the other observations.
- 198 The availability of a large number of remote-sensing systems at the observatory has enabled the possibility to both compare 199 and combine different techniques for studying atmospheric parameters (e.g., Mona et al., 2007; Madonna et al., 2010; Boselli 200 et al., 2012; Lopatin et al. 2013; Madonna et al., 2015). The synergy between lidars and photometer observations allows the 201 retrieval of vertical profiles of aerosol concentration for total, fine and coarse components through algorithms like GARRLiC 202 (Generalised Aerosol Retrieval from Radiometer and Lidar Combined data; Lopatin et al., 2013) and POLIPHON (Polarization 203 Lidar Photometer Networking; Mamouri and Ansmann, 2016; 2017). Ceilometers have shown good capabilities in the 204 detection of aerosol plumes even if with some limitations (Wiegner et al., 2014; Madonna et al., 2015). The combined use of 205 ceilometers and multiwavelength polarization Raman lidars can be an added value for acrosol variability investigation. 206 Additional cloud information provided by 24 h ceilometers can be precious for cloud masking prior to the analysis of lidar 207 measurements for aerosol profiling. These aspects are currently under investigation and developments within ACTRIS 208 implementation.
- 209 The combination of lidars and radars also demonstrated the enhancing power of synergistic observations: combination of lidar 210 and radar measurements during the Iceland volcanic eruption in 2010 showed radar capability of detecting giant volcanic
- 211 particles (Madonna et al., 2010; Madonna et al., 2013).
- Synergistic approach has been proposed for the study of thin liquid water clouds, combining multi-wavelength lidar and Doppler radar measurements (Rosoldi et al., 2022). It has been shown that microwave radiometer can be used to calibrate Raman lidar measurements for water vapour profiling and that the synergy between these instruments is an effective means for atmospheric water vapour monitoring (Madonna et al., 2006, Mona et al., 2007).
- However, despite its huge potential in atmospheric research, there are two major drawbacks associated with the remote sensing observations: the inability to conduct aerosol measurements under skies with low clouds and precipitations, along with the impossibility of characterising the particulate properties near the ground. Therefore, the recent implementation of the in-situ facility described in the next section is fundamental to achieve a complete characterization of the aerosol at the ground level where the aerosol particles directly affect ecosystems and human health. In addition, the in-situ measurements include the valuable chemical characterization of the particulate matter (PM), thus providing a deeper comprehension of the aerosol type,
- 222 the source apportionment and the mixing atmospheric processes.
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#### 224 **4 Description of the aerosol in-situ facility**

#### 225 **4.1 Overview**

226 The in-situ facility recently installed at CIAO comprises two main parts (Fig. 4): a field laboratory for aerosol particles online 227 measurements with continuous instrumentation and PM samplers and a chemical laboratory for the post-sampling analysis of 228 particulate aerosol particles collected over the on filters. This facility enables the measurement of all obligatory ACTRIS 229 aerosol in-situ variables: particle number concentration > 10 nm; particle number size distribution – mobility diameter 10 to 230 800 nm; particle light scattering & backscattering coefficient and particle light absorption coefficient and equivalent black 231 carbon concentration. Additionally, it allows the measurement of other four recommended variables: particle number size 232 distribution - aerodynamic diameter 0.8 to 10 µm; mass concentration of particulate organic and elemental carbon; mass 233 concentration of non-refractory particulate organics and inorganics and mass concentration of particulate elements.

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#### Figure 4: Outline and workflow of the <u>aerosol</u> in-situ facility at CIAO.

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The shelter has been designed according to the ACTRIS guidelines and recommendations (Wiedensohler et al., 2014), with the instrumentation arranged as follows: a dual spot aethalometer (AE33, Magee Scientific), a multi-wavelength integrating nephelometer (AURORA 3000, Ecotech) and an aerodynamic particle sizer (APS 3321, TSI) located <u>under\_downstream</u> a common  $PM_{10}$  (aerosol particles with an aerodynamic diameter less than 10 µm) inlet; a time-of-flight aerosol chemical speciation monitor (ToF-ACSM, Aerodyne Research), a scanning mobility particle sizer (SMPS,-3938<sub>2</sub>-TSI) and a condensation particle counter (CPC,-3750, -TSI) placed <u>under downstream</u> a  $PM_{2.5}$  (aerosol particles with an aerodynamic 245 diameter less 2.5 µm) common inlet. Additionally, two PMx samplers (SWAM 5a-Dual Channel Monitors, FAI Instruments) 246 and a PMx monitor (EDM 180, Grimm) are placed as standalone instruments with individual sampling lines. Additionally, 247 two PM<sub>x</sub> samplers (SWAM 5a-Dual Channel Monitors, FAI Instruments) are installed with respective inlets: one equipped 248 with two  $PM_{2.5}$  inlets, and the other with one  $PM_{10}$  and one  $PM_1$  inlet. Furthermore, a  $PM_x$  monitor (EDM 180, Grimm) is 249 placed as a standalone instrument with individual  $PM_{10}$  inlet line. Particular attention has been devoted to the design of the 250 common inlets and the sampling lines. The  $PM_{10}$  and  $PM_{2.5}$  common impactor type inlets, operating at a flow rate of 16.7 l 251 min<sup>-1</sup>, are compliant with EN 12341 and EN 14907 standards, respectively. The main challenge when transporting aerosol 252 particles to collectors and aerosol particles measuring instrumentation is to avoid aerosol particles-losses. Therefore, firstly, 253 the internal diameter of the main sampling pipe of the common  $PM_{10}$  and  $PM_{2.5}$  inlets must be such as to ensure that the sampled 254 air has a laminar flow along the entire path (Revnolds number less than 2000) to minimise the loss of particles by diffusion 255 and inertia. The instrument sublines (characterised by smaller inside diameters) are connected to the two common  $PM_{10}$  and 256  $PM_{25}$  inlets through their respective isokinetic flow splitter (Fig. 5), where the sample flow velocity closely matches the 257 velocity of the main flow. are connected to the common inlet through an isokinetic flow splitter where the sample flow velocity 258 is almost equal to the velocity of the main flow. Moreover, the ends of the tube in the isokinetic flow splitter must be sharp to 259 ensure a homogeneous distribution of the air sample to the instruments. Moreover, the tube ends in the isokinetic flow splitters 260 must be sharp to minimize turbulence and promote smooth airflow, ensuring uniform sampling. This design helps maintain 261 laminar flow, reduces aerosol losses, and enhances the accuracy and reliability of measurements. Another key feature of the splitter is that the sample is collected from the core of the main aerosol flow rather than from streamlines near the wall of the 262 263 main pipe, therefore, ensuring a representative sampling (especially for coarse and nanoparticles).



Figure 5: Image showing the rendering 3D	of the isokinetic splitters,	including the side v	iew and top view	(http://www.quattro-
esse.it/Home.html).				

The technical details of the common inlets and isokinetic splitters are shown in Table 12.

Common Inlet			Isokinetic splitter						
Inlet	Flow rate (1 min <sup>-1</sup> )	Int. Diameter (mm)	Speed (m s <sup>-1</sup> )	Reynolds Number	Instrument	Int. Diameter (mm)	Flow rate (1 min <sup>-1</sup> )	Reynolds Number	Speed (m s <sup>-1</sup> )
PM 10	16.7	21.2	0.8	1135	Aethalometer Nephelometer APS	8 8 4.4	<u>35</u> 5	885 885 320	1.6 1.6 1.09

PM 2.5	16.7	21.2	0.8	1135	SMPS CPC	4.4	2	655	2.2
					T <u>o</u> OF-ACSM	8	3	530	1

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Table 12: Technical details of the common inlets and isokinetic splitters.

272 All the sampling tubes are kept as short as possible and are placed in vertical position with bends and connectors avoided as 273 much as possible to suppress potential sources of turbulence, which would result in additional losses of particles. In addition, 274 the tubes are made of polyurethane antistatic material to guarantee perfect dissipation of accumulated static electricity, because 275 the static charges may remove significant portions of the aerosol to be sampled. In addition, in accordance with ACTRIS 276 recommendations, the tubes used are black sampling tubes supplied by TSI company (https://tsi.com/home/). These TSI 277 sampling tubes are made of conductive silicone, infused with carbon black to improve conductivity. This design is essential to 278 minimize electrostatic losses, which can occur in non-conductive tubes, such as those made of standard silicone or Teflon, 279 where particles can adhere to the tube walls due to static charges. The conductive nature of TSI tubes prevents the buildup of 280 electrostatic fields, thus improving particle penetration and reducing sampling distortions caused by particle loss.-The inlets 281 on the rooftop of the field laboratory are placed at one metre from each other and height of 1.5-2.0 m above the roof, 282 corresponding to approximately 4.5–5 m above ground level, with the aim of minimising local influences and potential 283 interferences in the sampling process.

In compliance with the ACTRIS indications, all the instruments in the laboratory are equipped with a Nafion dryer tube, a specialized device made from a sulfonated tetrafluoroethylene-based polymer. This device is used in aerosol sampling to remove water vapor from the gas stream while preserving the chemical integrity of aerosol particles (Monotube Dryer 700 (MD-700) - Perma Pure). These Nafion dryers-which keepsmaintain the RH well below 40%; under this threshold, in fact, changes in particle diameter due to RH variations are expected to be lower than 5%, thus obtaining comparable data, independent of the hygroscopic behaviour of the aerosol particles. Moreover, the upstream drying prevents the possible instrument damage caused by water condensation.

The Nafion dryers of Aethalometer, Nephelometer, APS and ACSM operate in a reflux mode, shown in Fig. <u>65</u>, which returns the dry sample back to the dryer for use as the purge after it has gone through the analyzer. Since this method uses all the dry sample as purge air, only the sample flow required for analysis passes through the dryer. This results in a high drying efficiency. <u>The vacuum on the purge air should be at least 15 inches Hg, with a higher vacuum preferable. This vacuum level is required</u> to provide the desired 2:1 purge to sample flow ratio based on the actual volumetric flow. <u>The vacuum on the purge air should</u> be at least 15 inches Hg, with a higher vacuum preferable. This vacuum level is required to provide the desired 2:1 purge-to sample flow ratio based on the actual volumetric flow. The 2:1 ratio ensures enough dry purge gas to continuously absorb
 moisture, preventing saturation and preserving sample integrity. This is crucial in aerosol particulate sampling, where even
 small amounts of moisture can affect particle characteristics and compromise measurement accuracy.

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#### Figure 65: Schematic Diagram - Nafion Reflux mode (MD-700-User-Manual, https://www.permapure.com).

Instead<u>In contrast</u>, the Nafion dryer connected to the SMPS and CPC, since the instruments need n butanol as a working liquid
 for the growth of aerosol particles, cannot operate in reflux mode but operates in counter flows using air dry coming from a
 compressor (Acoem 8301 LC-H Zero Air Generator). since the instruments need n-butanol as a working liquid for the growth
 of aerosol particles. -(Fig.ure 76).



**Figure <u>76</u>: Schematic Diagram - Nafion counter flows mode**.

- 309
- B10 Moreover, at the input\_inflow of each instrument, there is a high-resolution sensor connected to a software for continuous
- monitoring (every minute) of relative humidity and temperature. The accuracy of the sensors is 2.5% for the RH and 0.5°C for the temperature.

313 Lastly, in order to limit the temperature variation around the instruments, a continuously operating air conditioning system set

at 23-°C has been installed in the laboratory.

315 As previously mentioned, the in-situ facility is complemented by the chemical laboratory which enables complementary 316 measurements on the particulate matter-loaded filters coming from the PMx samplers, that is not possible to obtain with the 317 continuous instrumentation. The chemical laboratory include: an inductively coupled plasma optical emission 318 spectrophotometer (ICP-OES, series 5800, Agilent) used to perform the analysis of trace metals, and a multi-wavelength 319 OC/EC analyzer (DRI model 2015, Magee Scientific) used to analyse the carbonaceous fraction of the collected particulate. 320 The implementation of a such wide aerosol in situ measurements facility from scratch has required a total initial investment of 321 about 1M $\in$  and about 2 years, and about 100k $\in$ , 2 researchers with the support of a technicians are estimated to be needed to 322 operate the laboratory.

#### 323 **4.2 Instrumentation under the common PM<sub>10</sub> inlet**

324 As reported above, a  $PM_{10}$  common inlet is used to feed the aethalometer, the nephelometer and the APS. The aethalometer is 325 a key instrument for wildfire and pollution characterization, being able of detecting the fraction of particulate which absorbs 326 light, known as Black Carbon (BC), formed during the incomplete combustion of carbonaceous matter from biomass burning 327 and fossil fuel (Petzold et al., 2013). According to the ACTRIS guidelines, the AE33 aethalometer operating at seven different 328 wavelengths in the range 370-950 nm is used for the real-time monitoring of the concentration of BC. Briefly, the principle of 329 the aethalometer is to measure at given time intervals the attenuation of a light beam (at 880 nm) transmitted through a filter 330 where the particulate is continuously collected; the rate of change of optical transmission combined with the air flow rate 331 monitored through a mass flowmeter permits to determine the absorption coefficient, then converted into BC concentration by 332 means of the mass-absorption cross section. The dual spot technology refers to the contextual measurement of transmitted light 333 intensities through two separate spots of the filter at different loading levels, thus allowing to compensate for the so-called 334 loading effect largely described by Drinovec et al. (2015). The aethalometer is equipped with a sample stream dryer (Magee 335 Scientific) exploiting a semi-permeable Nafion membrane which keeps the RH well below 40%.

Among the other in-situ instruments placed under the  $PM_{10}$  inlet, the nephelometer can be considered in a certain way complementary to a ground-based lidar, expecting therefore to provide optical parameters consistent with those obtained from the lidar within the atmospheric planetary boundary layer (PBL). However, when the the PBL is particularly shallow (e.g., during wintertime), the nephelometer becomes the only tool to obtain the optical parameters of the aerosol<del>s particles</del> residing within the first hundreds of metres from the ground. The ACTRIS-compliant integrating nephelometer AURORA 3000 is used to measure the total scattering ( $\sigma_{sp}$ ) and the backscattering ( $\sigma_{bsp}$ ) coefficients (integrating within the angular range 9°-170° and 90°-170°, respectively), both correlated to the particle concentration (i.e., extensive properties). The peculiarity of the instrument is the utilisation of a light source emitting at three distinct wavelengths: the light at 635 nm (red) interacts strongly with large particulate matter such as desert dust and sea salt; the light at 525 nm (green) interacts strongly throughout the human range of visibility visible part of electromagnetic spectrum (smog, fog, haze); the light at 450 nm (blue) interacts strongly with fine and ultrafine particulates, such as wood fires and automobile combustion particulate. The nephelometer is equipped with a 36-inch-long Perma Pure Nafion MD-700 in order to prevent condensation of water droplets over the particles, which would increase their size and significantly change their scattering characteristics.

349 Lastly, the APS spectrometer provides high-resolution real-time aerodynamic measurements for the coarse fraction of the 350 particulate (Peters et al., 2003). The optical size range of the APS is from 0.37 to 20 µm, but since the spectrometer is connected 351 to a  $PM_{10}$  inlet and the counting efficiency of APS below 0.8  $\mu$ m aerodynamic diameter rapidly decreases and is unstable, the 352 realistically size range is from 0.8 to 10 um. The APS is based on the time-of-flight particle sizing, in which the aerodynamic 353 size of a particle determines its rate of acceleration, with larger particles accelerating more slowly due to increased inertia; the 354 time of flight between two laser beams is recorded and converted to aerodynamic diameter using a calibration curve. The 355 instrument measures in parallel the light scattering intensity of the sized particles in the equivalent optical size range from 0.8 356 to 10  $\mu$ m, thus providing further insights into the aerosol particles nature and composition.

The APS is connected to the sampling line just with the inner nozzle (sampling  $1 \ \text{l} \ \text{min}^{-1}$ ) from the common sampling line and the flow is dried by a 12-inch Perma Pure Nafion, while taking the additional sheath flow (4  $1 \ \text{min}^{-1}$ ) from the air compressor.

#### 359 **4.3 Instrumentation under the common PM2.5 inlet**

Even though the general ACTRIS recommendations for the in-situ measurements involve the analysis of the PM<sub>10</sub> fraction, the CPC, the SMPS and the ACSM represent an exception and are more conveniently placed under the cut-off size of a PM<sub>2.5</sub> inlet. The ACTRIS-compliant CPC is used to measure the number concentration of aerosol particles with diameter > 10 nm. In the CPC, an aerosol sample is continuously drawn through a heated saturator where the butanol is vaporized and diffused into the sample stream. Together, the aerosol sample and *n*-butanol vapour pass into a cooled condenser where the *n*-butanol vapour becomes supersaturated and condenses on the particle surface causing them to grow. The particles are then counted individually as they pass through a laser-based optical detector.

Regarding the SMPS, it is an instrument of interest for CIAO, being able to provide the size distribution and concentration of the fine fraction of the particulate in the size range 10 nm – 800 nm. It consists of four components in sequence: 1) a preimpactor which removes particles larger than the fixed upper limit of size; 2) a bipolar diffusion charger (model 3082, TSI) which confers a characteristic stationary charge distribution to the polydisperse particles by using a radioactive source (Kr-85); 3) a differential mobility analyzer column (DMA, model 3083, TSI) which separates the particles according to their electrical mobility by varying continuously the applied voltage within the column (Schmid et al., 2007); and 4) a condensation 373 particle counter (CPC, model 3750, TSI) where the classified monodisperse particles are counted after condensation of *n*-

butanol on their surface.

- The CPC and the SMPS are connected to the same 24-inch Perma Pure Nafion via a T-flow splitter in order to keep the RH below 40%. Moreover, a dry sheath air is needed for the SMPS to ensure particle sizing inside the DMA with a minimum fluctuation in RH and for this purpose a Silica Dryer Tube (model 3082, TSI) is incorporated in the DMA sheath flow system, which is a closed loop.
- 379 For what concerns the aerosol particles mass spectrometry techniques, the ToF-ACSM-(Aerodyne Research) has been shown 380 to be perfectly suited for the ACTRIS observatory platforms, It is specifically designed to provide continuous aerosol particle 381 monitoring over long time periods, spanning years, with the requirement of regular checks and calibrations to maintain the 382 accuracy and reliability of its long-term measurements. having been designed to provide continuous and unattended 383 measurements for aerosol monitoring on the timescale of years. The chemical speciation with high temporal resolution is a 384 unique feature of the ACSM technology, unobtainable with conventional filter sampling and subsequent post-processing 385 chemical methods; moreover, the ACSM is not subjected to sampling artefacts that affect the collection of semi-volatile PM 386 components by means of filters (Viana et al., 2006; Kim et al., 2015). The ToF-ACSM chosen for CIAO was introduced in 387 2013 (Fröhlich et al., 2013), providing a higher mass resolution (i.e.,  $m/\Delta m = 600$ ) and superior detection limits (i.e.,  $\frac{\langle ng m^3}{2}$ 388 less than nanograms per cubic meter) with respect to the previously developed quadrupole-ACSM (Ng et al., 2011) for a time 389 resolution of 30 min. The instrument measures the mass and chemical composition of non-refractory submicron aerosol 390 particles – i.e., organic substances, nitrates, sulphates, ammonium, and chloride – thus generating an invaluable database for 391 the research community to characterise the particulate sources and evolution. The operational principle of the instrument is 392 briefly described in the following: the aerosol enters the inlet where the aerodynamic lens efficiently samples and focuses 393 submicron particles to the subsequent vacuum chamber; here, the particles impact on a resistively heated porous tungsten 394 surface at approximately 600-°C which vaporises the non-refractory particulate; the vaporised matter is subsequently ionised 395 by electronic impact and detected through the ToF analyzer. In this case, the 24-inch Nafion dryer installed upstream the 396 instrument eliminates the complicating inlet effects due to particle composition dependent water uptake (Middlebrook et al., 397 2012).
- ACSM was installed in February 2023 and worked for some months in almost continuous way. Then some interventions were requested to accomplishing the optimization requests from ACTRIS aerosol in situ Ceentral Efacility, and the ACSM restarted operations just recently in April 2024. Anyhow the 3 months of almost continuous measurements performed in 2023 already provide some insights about aerosol <u>particles</u> present at the surface in Potenza. Fig<u>ure 87</u> reports daily concentrations for the 402 4 components as measured by ACMS in February-March-April 2023 period. Median values are preferred to mean ones for 403 avoiding the strong influence of outliers and spikes in the reported values. Monthly pie charts show the relevance of the

difference components for each one of the 3 months. As a general comment, we could say that the Potenza site is clearly a rural site with low PM concentration and a very high contribution of the organic substances. As a general comment, we could say that the Potenza site exhibits low PM concentrations and a very high contribution of the organic substances, as observed in rural areas-(see for comparison as example Atabakhsh1 et al., 2023 and Zhao et al., 2020). The observed increasing peaks in the total conentration but more pronounced in the organic component could be related to tree pollen events typically occurring in such period.









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Figure 87: Daily medians of the mass <u>concentration</u><u>fraction (a)</u> and mass <u>fraction</u><u>concentration (b)</u> of each of the 4 chemical components of non-refractory submicron aerosol particles observed at CIAO in February – March – April 2023.

#### 415 **4.4 PMx samplers and PMx monitor**

416 Additionally to the online instruments report above, the field laboratory is equipped with two PMx samplers for the continuous 417 sampling and concentration measurement of PM<sub>10</sub>, PM<sub>2.5</sub> and PM<sub>1</sub> (aerosol particles with an aerodynamic diameter less than 1 418 µm) mass fractions collected over both Teflon and quartz filters; the determination of the mass of collected samples is based 419 on the  $\beta$ -ray attenuation equivalent method, which strongly reduces the workload and the operator-associated variability if 420 compared to the standard gravimetric method (Baltensperger et al., 2001). In particular, the device measures the attenuation 421 of  $\beta$ -ray across the filter medium which collects particulate matter, and the attenuation of intensity in  $\beta$ -ray is proportional to 422 the amount of material present. Each PMx sampler is equipped with two independent sampling lines (i.e., PM<sub>10</sub>/PM<sub>2.5</sub> and 423 PM<sub>2.5</sub>/PM<sub>1</sub>), thus enabling the simultaneous collection of different PM fractions on independent filters. According to the 424 workflow reported in Fig. 4, the particulate collected over the filters is subjected to further analysis within the chemical 425 laboratory: the PM<sub>10</sub>, PM<sub>2.5</sub> and PM<sub>1</sub> collected over 24h on Teflon filters are analysed to determine the concentration of metals

- 426 by means of the ICP-OES. On the other hand, the PM<sub>2.5</sub> collected over 24h on quartz fibre filters are analysed to quantify the 427 organic carbon (OC) and elemental carbon (EC) fractions using the thermal optical method by the OC/EC analyzer; the 428 utilisation of quartz fibre filters for the OC/EC analysis is strictly recommended by the WMO/GAW 2016 guidelines, and it 429 constitutes the only exception to the Teflon filters commonly used for other analyses. In fact, the particulate collected on Teflon 430 filters is not limited to ICP-OES analysis but can also be analyzed through alternative techniques such as X-ray fluorescence 431 (XRF) and Particle Induced X-ray Emission (PIXE) in order to find complementarities between the three techniques for the 432 determination of a range of metals. 433 Furthermore, even if not included in the mandatory ACTRIS variables to be measured, the mass concentration for the cut-off
- 434 diameters of  $PM_{10}$ ,  $PM_{2.5}$  and  $PM_1$  belongs to the set of standard measurements to monitor the particulate matter, providing 435 insight into the separation of fine and coarse particles within the aerosol.
- The PMx monitor operating at CIAO currently represents one of the main automated measurement systems for studying the concentration levels of particulate matter in ambient air. Based on the detection principle of the light scattering at the level of single particles, the system offers simultaneous real-time measurements of  $PM_{10}$ ,  $PM_{2.5}$  and  $PM_1$  and particle number distribution with a resolution of 0.1 µg m<sup>-3</sup>.

#### 440 **4.5 Chemical laboratory**

441 The CIAO chemical laboratory is equipped with an ICP-OES and an OC/EC analyzer. The ICP-OES (5800 series, Agilent) is 442 used to determine the qualitative and quantitative elemental composition of the metals present in the atmospheric particulate with high sensitivity, at values below the 1  $\mu$ g l<sup>-1</sup> limit for certain elements. The metals are introduced into the atmosphere 443 444 from various anthropogenic and natural sources. Anthropogenic metals are released into the atmosphere during combustion of 445 fossil fuels and wood, as well as during high temperature industrial processes and waste incineration; natural emissions result 446 from a variety of processes acting on crustal minerals, including volcanism, erosion, surface winds, forest fires and ocean 447 evaporation (Allen et al., 2001; Pakkanen et al., 2001; Rajšić et al., 2008). Various metals are used as marker for the 448 identification of emission sources: aluminium and silicon are primarily derived from soil and rocks (crustal elements); sodium 449 and chlorine are typically associated to marine aerosols particles; arsenic, cadmium, manganese and lead mostly derive from 450 combustion of fossil fuels occurring at high temperature, to name a few. The ICP-OES analysis of particulate matter requires 451 a preliminary microwave digestion of the filter in acidic conditions to extract the metals, carried out by means of a microwave 452 digester (ETHOS UP, FKV). The obtained liquid sample is then nebulized and introduced into the plasma as an aerosol 453 suspended in the argon gas: due to the high temperatures within the plasma (7000 - 10000 K), a significant fraction of most 454 elements exists as atoms or ions in the excited state, causing an intense polychromatic emission which continuously brings 455 back the elements to their ground state. The polychromatic emitted light is dispersed into individual wavelengths by a 456 polychromator and detected by a photosensitive charge-coupled device (CCD). The concentration of each metal in the sample is obtained by using a calibration curve referred to a solution containing the analysed elements of known concentration.

458 The multi-wavelength OC/EC analyzer (2015 DRI, Magee Scientific) compliant with ACTRIS is used to quantify the total 459 carbonaceous content of the particulate matter (total carbon, TC) and the OC and EC subfractions. EC is essentially a primary 460 pollutant, emitted directly from the incomplete combustion of fossil fuels and the pyrolysis of biological material during 461 combustion, whereas OC can be directly emitted from the incomplete combustion of organic materials and the degradation of 462 carbon containing products such as vegetation - primary OC - or produced from atmospheric reactions, involving gaseous 463 organic precursors, i.e., secondary OC (Zhou et al., 2006). The operational principle of the thermal/optical analysis is based 464 on the preferential desorption of OC and EC materials under different temperatures and atmospheres programmed within 465 specific thermal protocol, such as the EUSAAR 2 (Cavalli et al., 2010) which is currently used within the ACTRIS community. 466 OC usually desorbs under a non-oxidising helium atmosphere at temperatures up to 570-°C, while the EC is combusted in an 467 oxidising atmosphere with 2% O<sub>2</sub> at temperatures up to 850-°C. However, since part of the OC turns into the light-absorbing 468 pyrolytic carbon which desorbs during the oxidising mode, the correct discrimination between the OC and the EC fractions is 469 conveniently identified with the point at which the light transmission reaches the pre-pyrolysis value. The liberated carbon is 470 then completely oxidised to carbon dioxide passing through a heated catalyst MnO<sub>2</sub> and finally quantified by an NDIR (Non-471 Dispersive Infrared) detector.

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#### 473 5 Synergistic Combined deployment of aerosol remote sensing and in-situ measurements

474 Synergistic approaches Combined approaches combiningusing aerosol particles profiling and in-situ measurements are one of 475 the most beneficial strategies in aerosol research, allowing an accurate typing and estimation of the impacts of particulate 476 matter (Molero et al., 2020). Remote sensing techniques provide the vertical profile of the particle size distribution of the 477 aerosol particles as well as further physical and optical properties useful for understanding complex atmospheric phenomena 478 (Vratolis et al., 2020); however, they are not able to provide information under cloudy sky conditions or at the ground level, 479 where the identification of aerosol <u>particles</u> type is only possible using the in-situ instrumentation. The in-depth typing of the 480 aerosols particles require the information on the chemical composition, attainable only by means of in-situ measurements. The 481 complete set of data resulting from the combined approaches is crucial for identifying the sources and the evolution of 482 concentration levels of particulate matter over time (Bressi et al., 2021), and it is of paramount importance for the 483 implementation of controls or policies to reduce aerosols that negatively affect air quality and public health.

484 The complete picture of the aerosol particles-typing is also expected to clarify further the climate effects of particulate matter. 485 In fact, the estimation of the radiative effect of atmospheric aerosol particles requires the knowledge of multiple parameters, including the aerosol <u>load\_concentration</u>, the optical properties, the chemical composition, the presence of clouds and the albedo of the underlying surface. The accurate identification of aerosol particle types is also needed to improve the understanding of atmospheric dynamics and long-range transport, to improve satellite aerosol retrieval algorithms, and to validate climate models.

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491 The multiwavelength polarisation Raman lidar is a well-established active remote sensing technique for the detection and 492 characterization of aerosol-types (Nicolae et al., 2018; Papagiannopoulos et al., 2018). Specifically, it can provide vertically 493 resolved information on extensive (e.g., aerosol backscatter coefficient, aerosol extinction coefficient and volume 494 depolarization ratio) and intensive (e.g., Ångström exponent, lidar ratio and particle depolarization ratio) aerosol optical 495 properties. The extensive properties depend on the aerosol particles concentration, whilst intensive ones are type-sensitive 496 providing indication about the particle size, shape, and indices of refraction that allow for the characterization of different 497 aerosol types. Nevertheless, the intensive properties might not be sufficient to guarantee accurate typing, as some aerosol types 498 (e.g., volcanic and desert dust particles) have very similar intensive properties but are attributed to different sources and 499 generating mechanisms. For this reason, the discrimination of aerosol particles that typically have the same optical 500 characteristics calls for the combined use of lidar observations and transport model simulations.

- Finally, <u>T</u>the aerosol in-situ observations can help in the assessment of the uncertainty of remote sensing-retrieved products
   like mass concentration, refractive index and fine-particle concentration obtained through inversion algorithms (e.g.,
   Veselovskii et al., 2012; Lopatin et al., 2013).
- 504 <u>Furthermore, the availability of collocated in situ and remote sensing measurements of aerosols also represents an added value</u>
- 505 for modelling. Indeed, it can contribute to the increase in the accuracy of model predictions, allowing the reduction of the
- 506 uncertainty of aerosol measurements in the atmosphere (e.g., Vratolis et al., 2020), as well as to a better evaluation of aerosol
- 507 models. In recent years, collocated datasets have also been increasingly utilized for training machine learning-based models,
- 508 as demonstrated by Redemann and Gao (2024).
- 509 To emphasize the potential synergy and added value of combining in-situ and remote sensing techniques, Table 2 presents a
- 510 <u>comprehensive list of instruments and the respective parameters they measure.</u>

	Instruments	Parameters
	Ka-band Doppler radar Metek MIRA-35	Vertical profiles of Doppler spectrum moments (e.g. Signal-to-noise ratio, Doppler velocity, spectral width, equivalent reflectivity factor) and of linear depolarization ratio of atmospheric targets (clouds, precipitations, insects, giant aerosols). Typical measurement range: 100m-15km
	Compact Ka-band Doppler radar MIRA 35C	Same as above. Typical measurement range: 100m-12km
	W-band Doppler radar RPG-FMCW-94	Same as above. Typical measurement range: 50m-12km
Cloud Remote Sensing	K-band Doppler radar Metek MRR-PRO	<u>Vertical profiles of hydrometeors' Doppler spectrum</u> <u>moments, of raindrop size distribution, liquid water mass</u> <u>concentration and rain rate. Typical measurement range:</u> <u>15m-6km</u>
	Microwave radiometer RPG-HATPRO-G5	<u>Vertical profiles of temperature, humidity and cloud liquid</u> <u>water mass concentration; integrated precipitable water</u> <u>vapor and cloud liquid water path. Typical measurement</u> <u>range: 0-10km</u>
	<u>Ceilometer Vaisala CL51</u>	Vertical profiles of attenuated backscatter coefficient from atmospheric particles (aerosols, clouds and precipitations), cloud base heights, aerosol vertical layering. Typical measurement range: from near surface (15m) up to 12km for clouds and depending on the aerosol load in a cloud- free atmosphere.
	<u>Ceilometer Vaisala CL31</u>	Same as above. Typical measurement range: from near surface (15m) up to 8km for clouds and depending on the aerosol load in a cloud-free atmosphere.
	Ceilometer Lufft CHM15k	Same as above. Typical measurement range: from near surface (15m) up to 15km for clouds and depending on the aerosol load in a cloud-free atmosphere.
	2 Doppler lidars Halo Photonics Stream LineXR	Vertical profiles of attenuated backscatter coefficient from atmospheric particles (aerosols, clouds and precipitations) and of vertical/horizontal wind components.

<b>Trace gases Remote Sensing</b>	FTIR Bruker 125HR	<u>Infrared spectrum contain the signatures of vibrational-rotational transitions of numerous trace gases in the terrestrial atmosphere as they absorb solar radiation (O<sub>3</sub>, <u>HCl, HF, ClONO<sub>2</sub>, HNO<sub>3</sub>, N<sub>2</sub>O, CH<sub>4</sub>, CO, C<sub>2</sub>H<sub>6</sub>, and <u>HCN).</u></u></u>
	Fixed multi-wavelength Raman lidar	Aerosol extinction coefficients at 355 and 532 nm; Aerosol backscatter, Volume and particle depolarization ratio at 355, 532 and 1064 nm and water vapour mixing ratio
	Mobile multi-wavelength Raman lidar	Aerosol extinction coefficients at 355 and 532 nm; Aerosol backscatter, Volume and particle depolarization ratio at 355, 532 and 1064 nm
nsing	MUSA Transportable Fixed multi-wavelength Raman lidar	Aerosol backscatter at 355, 532 and 1064, Aerosol extinction coefficients at 355 and 532 nm, Volume and particle depolarization ratio at 532
te Sei	Scanning UV Raman lidar	Aerosol backscatter, Aerosol extinction coefficients, Volume and particle depolarization ratio at 355
Aerosol Remo	Automatic sun/sky/lunar photometer Cimel 318T	<u>Aerosol Optical Depth (AOD), Volume Size Distribution</u> (VSD), complex refractive index (n), shape factor, water vapor content.

	<u>SMPS</u>	Size distribution and concentration of the particles in the size range 10 – 800 nm
	APS	$\frac{\text{Size distribution and concentration of the particles in the}}{\text{size range } 0.8 - 10 \mu\text{m}}$
	<u>CPC</u>	$\frac{\text{Concentration of the number of particles with dimensions}}{\geq 10 \text{nm}}$
situ	<u>Nephelometer</u>	Aerosol scattering and backscattering coefficients at 450, 525 and 635 nm
ol in-	Aethalometer	Aerosol absorption coefficients and BC concentration at seven wavelengths in the range of 370-950 nm
Aeros	<u>PM<sub>x</sub> Monitor</u>	Mass concentration for the cut-off diameters of PM <sub>10</sub> , PM <sub>2.5</sub> and PM <sub>1</sub>
	ACSM	Real-time chemical characterization of the main organic and inorganic components of the non-refractory sub- micrometric aerosol particles
	<u>ICP-OES</u>	Qualitative and quantitative elemental composition of the metals on collected filters of aerosol particles
	OC/EC Analyzer	TC and subfraction OC/EC concentrations on collected filters of aerosol particles

## 511 512

Table 2: List of all CIAO remote sensing and in-situ instruments and respective parameters.

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514 In the following subsections we present three emblematic cases recurring at CIAO where the combined deployment of the in-515 situ and remote sensing observations is expected to be of added value: 1) Wildfires become more and more relevant in the 516 Mediterranean, especially in view of the changing climate that is expected to increase temperature and in turn will affect their 517 frequency, duration and intensity in the next decades. In this context, small and local fires are widely distributed distributed, 518 and their characteristics and assessment could be important at global level. De Rosa et al. (2022) showed with the use of lidar 519 observations that fresh fires can be surprisingly characterised by low absorption; this would imply a different impact of local 520 fires in the radiation budget which requires investigation and validation by means of in-situ measurements. 2) Local pollution 521 during winter and adverse weather can be investigated in a more exhaustive manner only by in-situ observations, since lidar 522 observations provide very little information due to the generally low and unresolved by lidars PBL height. 3) Desert dust 523 intrusions often reach Europe and especially the Mediterranean Basin affecting local air quality, health and ecosystem and

524 socio-economic sectors (e.g., Monteiro et al., 2022). Given all the above, the deployment of in-situ measurements at well-525 equipped sites like CIAO is crucial to quantify the impact at the ground level.

#### 526 5.1 Local wildfires

The study of smokes from wildfires spreading <u>in\_over</u> short distance represents a great example for a synergistic approach based on remote-sensing and in-situ techniques. In such a case, the smoke particles spread mainly at low levels and deposit fast on the ground, where in-situ measurements are the only tool to provide reliable information to support and integrate what is observed above medium overlap region, a prerogative of remote-sensing techniques.

- 531 The multiwavelength polarisation Raman lidar is a well-known tool to study smoke layers in the atmosphere, being able to 532 separate aerosols according to their specific optical signature (Ohneiser et al., 2021). Specifically, a sign of the dominance of 533 smoke in the aerosol laver is the aerosol extinction-to-backscatter ratio (the so-called lidar ratio, S) at 532 and 355 nm, which 534 is typically high (i.e., > 50 sr) as a consequence of the presence of absorbing BC produced during the biomass burning; 535 moreover, the ratio of S measured at the different wavelengths may be used as an indicator of the phase of the ongoing wildfire 536 (e.g., Nicolae et al., 2013). Other lidar parameters largely used to investigate the smoke are the particle linear depolarization 537 ratio (PLDR) and the Ångström exponent (AE), which provide information about the shape and the size of the particles, respectively. In the case of a local wildfire, the observation of quasi-spherical and relatively small particles is expected, since 538 539 the newly produced smoke particles do not have the time to undergo modifications during transport.
- 540 The Ångström absorption and scattering exponents (AAE and SAE) derived from the aethalometer and nephelometer 541 measurements, respectively - provide the optical typing of the smoke, with the value of AAE expected to correlate with the 542 lidar observations (Cazorla et al., 2013) and, therefore, to the nature of spreading fire.
- 543 Among the aerosol in-situ instruments, the aethalometer is crucial to study smokes produced during wildfires, being able to 544 quantify the BC originated from the incomplete combustion of carbonaceous matter and providing an estimate of the biomass 545 burning (BB) apportionment to the overall BC (Sandradewi et al., 2008). Furthermore, particles resulting from the incomplete 546 combustion have been reported to contain a significant organic carbon fraction, including numerous known toxic and 547 earcinogenic polycyclic aromatic hydrocarbons (PAHs) (Nelson et al., 2021). To access relative amounts of organic carbon, 548 the method involves comparing two optical indicators of carbonaceous particulate matter derived from the aethalometer 549 measurements: BC at 880 nm, measuring elemental carbon that absorbs a broad spectrum of wavelengths, and UVPM at 370 550 nm, measuring particulate matter that, due to increased organic carbon content, absorbs disproportionately in the UV range 551 compared to BC (Olson et al., 2015), UVPM, also known as brown carbon, is associated with toxic species such as PAHs and 552 has been observed to be elevated in smoke resulting from biomass burning (Huangh et al., 2018). Additionally, the OC/EC 553 thermal/optical analysis on  $PM_{2.5}$  fraction is very important because the increase of organic carbon and elemental carbon

554 concentrations has been the most indicated as an element that reflects wildfire emissions. The fine particles, particles generally 555 2.5 um in diameter or smaller, represent a main pollutant emitted from wildfire smoke so other important in situ analyses of 556 the travelled smoke are the size distribution and concentration of fine particles by the SMPS and CPC, respectively. In fact, it 557 is expected that the fine mode will be more densely populated and concentrated during these events compared to the rest of 558 the year. Further confirmation of the increase in fine and ultrafine particulate matter during fire events is given by the 559 PM2.5/PM10 and PM1/PM2.5 ratios obtained from the real-time measurement of PM10, PM2.5 and PM1 concentrations using the PMx monitor. In fact, the mean fraction of fine PM (PM25/PM10) and ultrafine PM (PM1/PM25) is expected to be significantly 560 561 higher during the fire period compared to the non-fire period. Fine particles ( $\leq 2.5 \mu m$ ) are a major pollutant from wildfire 562 smoke. Key in-situ analyses include size distribution and concentration measurements using SMPS and CPC, as fine particles 563 are more abundant during fires compared to other periods. Real-time  $PM_x$  monitoring confirms increases in  $PM_2$  5/ $PM_{10}$  and 564  $PM_1/PM_{2.5}$  ratios during fire events. Finally, the in-situ investigation of wildfire smoke is completed by the chemical analysis 565 obtained with the ToF-ACSM: in particular, key tracers of biomass burning organic aerosol particles in mass spectra are the enhanced signals at m/z 60 and 73 attributed to  $C_2H_4O_2^+$  and  $C_3H_5O_2^+$  ions, respectively, coming from the fragmentation of 566 567 the so-called "levoglucosan-like" species originated from the pyrolysis of cellulose (Cubison et al., 2011). Finally, the chemical analysis of the filters through the ICP-OES is fundamental for tracking the levels of potentially toxic elements (PTEs) such as 568 569 As, Sb, Cd, Hg, Pb, Cr, Cu, Ni, Se, Tl, Sn, V, and Zn. This monitoring is vital as these elements have the potential to be 570 released into the environment during wildfires, posing a threat to humans and animals when their absorbed doses surpass the 571 established reference values (Pacifico et al., 2023).

572 <u>This case underlines the critical need for a combined approach, where in-situ measurements bridge the gap and enhance the</u>
 573 <u>interpretation of remote-sensing data, showcasing the strength of CIAO's integrated monitoring capabilities.</u>

#### 574 **5.2 Local pollution in wintertime**

Winter months commonly exhibit heightened air pollution levels, primarily attributed to temperature inversions. Inversion occurrences involve a layer of warm air confining colder air and pollutants close to the ground, impeding their dispersion into the atmosphere. Unlike summer air pollution, winter conditions result in the prolonged presence of pollutants, increasing the likelihood of higher inhalation rates. This extended exposure raises health concerns for individuals, as reduced ventilation and dispersion contribute to potential health effects.

Air quality near the ground during winter is expected to be dominated by local residential heating emissions with the contribution of vehicle engine exhausts. For this season, the in-situ measurements represent the most viable way to investigate the aerosol <u>particles</u> distribution and composition, while the deployment of remote sensing instruments (e.g., lidar) is limited by instrumental and environmental factors. During wintertime, the condensation of water droplets (especially during nighttime) along with the recurrent formation of cloud layers attenuate the laser beam, thus impeding the lidar/ceilometer measurementsretrievals; moreover, even under clear sky conditions, the particulate is usually confined within the first 300 m from the ground (i.e., the typical PBL layer thickness in wintertime), where the active remote sensing techniques are not able to provide reliable results, because of the typically not complete overlap between laser beam and receiving system in lidar at this vertical range.

- 589 The climatological profile of aerosol backscatter at 532 nm for winter season at CIAO ((blue line in Fig. 9) shows very clean
- 590 air respect to other seasons in the whole investigated atmospheric column (https://doi.org/10.57837/cnr-imaa/ares/actris-
- 591 <u>earlinet/level3/climatological/2000\_2019/pot).</u>



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Figure 9: Climatological profiles of the aerosol backscatter at 532 nm observed at CIAO from 2000 to 2019. Normal
 Season average is reported: each profile is the average of all aerosol backscatter profiles observed at CIAO for each
 season in routinely measurements collected from 2000 to 2019.

596

597 Only the last point close to the surface is slightly higher respect to the other seasons, but the information content is too low for

598 <u>further investigation. These cases are typically considered as clean day from the aerosol remote sensing perspective. However,</u>

- 599 it must be noted that lidar is blind in the lowest portion of the atmosphere, where pollutants are concentrated due to the low
- 600 <u>boundary layer height (PBL).</u>

601 On the other hand, only the in-situ measurements, which do not see above the boundary layer height, can well capture what's

602 <u>happening close the surface.</u>

the in-situ instrumentations enable the analysis of the particulate matter collected at the ground level where the pollutants highly concentrate as a result of the stagnant, dense and cold air. Among the in-situ measurements, aethalometer plays a key role: the BC content of particulate matter originates mostly from the incomplete combustion of both fossil fuel and biomass used as combustibles for domestic heating. The BC content is expected to be higher with respect to the background summer levels, especially due to the contribution of local residential heating and the air stagnation. The BC source apportionment is expected to be determined by both biomass burning fraction due to the residential wood burning and the fossil fuel due to the traffic exhaust and residential heating.

610 The OC/EC analysis on the PM<sub>25</sub> fraction is expected to provide additional data to both support and integrate the results 611 obtained with the aethalometer (Schmidl et al., 2008; Gonealves et al., 2010; Pio et al., 2011; Sirignano et al., 2019). The 612 nephelometer is expected to provide the optical parameters at the ground complementing the ones obtained by lidar 613 measurements in the 300 m UTLS region; the total scattering coefficient  $\sigma_{sb}$  and the backscattering coefficient  $\sigma_{bb}$  are related 614 to the concentration of particles, with a dominant response expected at 450 nm and relatively high values for the SAE, 615 corresponding to the fine and ultra fine particles typically produced by heating emissions (Esteve et al., 2012). Further 616 elucidations on the nature and the origin of the particulate can be certainly obtained with the ToF ACSM. In this case, an 617 accurate prediction of the chemical composition of the particulate is not a trivial task since many factors contribute to the 618 chemistry of the particulate and, as of today, there are no previous data reports for such type of analysis at the site in wintertime. 619 In principle, however, the chemical speciation of the PM<sub>+</sub> fraction from the ACSM is expected to put in evidence a prevalence 620 of the organic matter derived from the combustion processes. Moreover, as previously reported (Chen et al., 2012), during 621 wintertime the recurrent exceedances of the fine particle fractions may be due to the abundance of the secondary ammonium 622 nitrate ( $NH_4NO_4$ ), attributed to residential wood combustion and diesel engines through the emission of nitrogen oxides ( $NO_4$ ) 623 from these sources. Finally, the importance of the chemical analysis of the filters must be underlined; through ICP OES the 624 main metals present in the particulate can be analyzed (Na, Mg, Al, Ca, V, Cr, Fe, Mn, Ni, Cu, Zn, As, Mo, Sb, Cd, Ba, Pb) 625 which come from specific sources, such as the combustion of fossil fuels in industries or power plants or in vehicle combustion 626 engines, coal and wood combustion processes, non combustion related emissions from vehicular traffic and dust resuspension 627 phenomena resulting from traffic (Dušan et al., 2017; Zhi et al., 2021).

In th<u>is context</u>,<u>e following</u> we investigate the average daily concentration of elemental carbon (eBC) obtained by the aethalometer (Figure <u>10</u>8a), covering the period from June 2023 to April 2024, to have a first insights into air quality near the ground during winter. Our analysis reveals no significant increase in eBC concentration during the winter months compared to background levels observed in summer. However, when examining the daily average percentage of black carbon (BC) originating from biomass burning (BB%), as determined by the Sandradewi model (Sandradewi et al., 2008), in conjunction with the daily average temperature data obtained from the Vaisala AWS310 weather station situated at the site (Figure <u>108b</u>), an intriguing trend emerges. It becomes evident that BB% is substantially higher during winter months than during summer months. Given the minimal occurrence of wildfires and prescribed burns at the site during winter, the primary source of biomass burning influence can be attributed to residential burning, a consequence of the notably low temperatures experienced during that period.



638 639

Figure <u>108</u>: Daily average eBC concentration obtained by aethalometer from June 2023 to April 2024 a); daily average BB% determined by the Sandradewi model and daily average temperature data obtained from the Vaisala AWS310 weather station from June 2023 to April 2024 b).

<sup>644</sup> These first data indicate that the main source of BC during winter at our site is predominantly from local residential heating 645 emissions. Unlike remote sensing, which may suggest clean air conditions, in-situ measurements reveal significant pollution

- 646 <u>at ground level, underscoring that winter cases should not be considered background conditions in the the boundary layer</u>
- 647 region. Therefore, this preliminary study demonstrates that combining remote sensing and in-situ measurements provides
- 648 <u>critical insights that neither method can achieve independently.</u>
- 649 <u>Future studies could expand on this by incorporating additional in-situ instruments alongside the aethalometer.</u>
- 50 For example, OC/EC analysis on the PM<sub>2.5</sub> fraction could support and complement aethalometer results (Schmidl et al., 2008;
- 651 Gonçalves et al., 2010; Pio et al., 2011; Sirignano et al., 2019), while a nephelometer could provide ground-level optical
- parameters, such as the scattering coefficient ( $\sigma_{sp}$ ) and backscattering coefficient ( $\sigma_{bsp}$ ), which are linked to particle
- 653 concentrations, particularly fine and ultrafine particles typical of heating emissions (Esteve et al., 2012).
- 54 <u>The ToF-ACSM could offer a detailed chemical composition of PM<sub>1</sub>, expected to reveal a dominance of organic matter from</u>
- 655 combustion processes. Secondary ammonium nitrate (NH4NO3), associated with residential wood burning and diesel emissions
- (via NOx), could also explain exceedances in fine particle fractions during winter (Chen et al., 2012).
- 657 Finally, filter analysis with ICP-OES could identify metals in the particulate matter (e.g., Na, Mg, Zn, Pb) originating from
- 658 specific sources such as fossil fuel and biomass combustion, vehicular traffic, and dust resuspension (Dušan et al., 2017; Zhi
- et al., 2021). This in-situ multi-instrument approach would provide a comprehensive view of particulate composition and
- 660 sources during winter pollution episodes at the surface, enabling a deeper insight into winter aerosol conditions, addressing air
- 661 <u>quality challenges, and accurately evaluating their health impacts.</u>

#### 662 **5.3 Dust intrusions**

- During summer and spring, the site is regularly affected by Saharan dust intrusions (Mona et al., 2006). Desert dust particles
  have many effects: they can impact climate, the precipitation cycle, and human health (Sokolik et al., 2007; Mona et al., 2023).
  Mineral dust particles can act as cloud condensation nuclei (CCN) and thereby determine the concentration of the initial
- droplets, albedo, precipitation formation, and lifetime of clouds (Levin et al., 1996; Levin et al., 2005).
- 667 The multiwavelength polarisation Raman lidar provides highly resolved spatial and temporal atmospheric profiles that allow
- for the separation of the different aerosol layers (Pappalardo et al., 2004b; Papagiannopoulos et al., 2018). Large and irregular
- shaped Saharan desert dust particles produce medium lidar ratios S, relatively high PLDR values and they are spectrally neutral
- to backscatter and extinction producing low Ångström exponent referred to the wavelengths 355-532 nm (Freudenthaler et al.,
- 671 2006; Fernandez et al., 2019). In fact, mineral desert dust aerosols particles predominantly consist of coarse mode particles of
- 672 irregular shapes (Mahowald et al., 2014).
- 673 In-situ measurements, in case of sedimentation events, provide complementary information on the advected dust. Low values
- of nephelometer-derived SAE that indicate coarse particles and, conversely, high aethalometer-derived AAE values that
- demonstrate the wavelength dependent absorption (Cazorla et al., 2013).

- 676 When the atmosphere is dominated by particles with large dimension such as dust particle, the sedimentation is fostered and 677 involves higher return to the ground level so the measurements of size distribution of coarse particles by APS plays a key role 678 in dust studies Furthermore, the low PM2.5/PM10 ratio obtained by real time measurements using the PMx monitor could be 679 the confirmation that the main component of the desert dust events is the PM coarse fraction. Finally, the PM<sub>10</sub> mass 680 concentration collected over 24h on filter measured by the PMx sampler (SWAM 5a-Dual Channel Monitors, FAI Instruments) 681 will be higher during a dust event compared to non-dust events, with PM10 concentration values that could exceed the European 682 daily limit value (2008/50/CE European directive). 683 Regarding the chemical characterization, the ICP OES plays a key role in evaluating the influence of the transport of dust by
- detecting the elemental composition of the mineral fraction. In particular, monitoring the concentrations of the typical crustal
   elements such as As, Al, Ca Cr, Cd, Co, Cu, Fe, K, Mn, Mo, Na, Ni, Pb, Sb, Se, Sn, and Zn and Rare Earth Elements (REEs)
   is relevant because are generally markedly higher during desert dust event than in comparison with their annual means (Aydin
   et al., 2012: Rodriguez Navarro et al., 2018: Mărmureanu et al., 2019).
- 688 During dust-dominated atmospheric conditions, sedimentation increases, returning large particles to the ground. The APS size
- $\frac{\text{distribution measurements of coarse particles are crucial for dust studies. A low PM_{2.5}/PM_{10} ratio from real-time PMx monitor}{\text{M}_{2.5}/PM_{10}}$
- $\frac{data \text{ confirms the dominance of the coarse fraction during desert dust events. Additionally, 24-hour PM_{10} mass concentrations}{measured by the PM_x sampler (SWAM 5a-Dual Channel Monitors, FAI Instruments) are higher during dust events, often$
- 692 exceeding the European daily limit (2008/50/CE European directive).
- 693 For chemical characterization, ICP-OES detects the mineral fraction's elemental composition, particularly crustal elements
- 694 like Al, Ca, Fe, K, and Na, along with Rare Earth Elements (REEs). These concentrations are significantly elevated during
- dust events compared to annual averages (Aydin et al., 2012; Rodriguez-Navarro et al., 2018; Mărmureanu et al., 2019).
- 696

697 In the following we report an example of aerosol remote sensing and in-situ observation for a Saharan dust intrusion at CIAO 698 to demonstrate the possibilities for synergistic complementary combination of data from lidar and in-situ aerosol measurements. 699 Even if only the APS instrument was available at that time, the presence of just one in-situ instrumentation already shown the 700 importance of such combination of techniques. The observations are related to the second half of June. Figure 9a reports the 701 fine mode fraction as retrieved from CIAO photometer measurements and available at aeronet.gsfc.nasa.gov. This parameter 702 provides information about the fraction of fine mode particles respect to the coarse one as obtained from the AOD (Aerosol 703 optical Depth) measurements. This parameter is retrieved from columnar measurements and therefore refer to the total 704 atmospheric column. Fig 9a clearly shows that in the 20-23 June period the coarse particles are more abundant respect to 705 previous and following period. For the same period Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) 706 backward trajectories ending over Potenza indicate Sahara desert as potential source of the observed particles.

Lidar obseravtionsobservations provide a better insight of the temporal and vertical distribution of the aerosol <u>particles</u> at CIAO on those days. Figure 9b-c report available Lidar observations for the period. It shows the color-maps of the vertical distribution and temporal dynamics of the aerosol as time series of range-corrected lidar signals at 532 nm for the night of 22 June 2023 and the daytime day of 23 June 2023. In particular, these plots report the component of backscattered signals at 532 cross-polarized respect to the emitted laser light: the presence of high cross-polarized backscatter signals is a signature of presence in that portion of 4d atmospheric region of aspherical particles, like Saharan dust ones.

The representation of the aerosol\_particles distribution during the night of 22 June (Fig. 9b) shows two main layers of dust: one at an altitude close to 1 km above ground level (agl) and a second denser one above it at a height of approximately 3 km agl. Particularly interesting for the potential link with in-situ measurements is a branch of the lower layer around 01:30 in the night between 22 and 23 June, which seems to descent in altitude and could potentially sediment at the ground. It is worth to note that the lidar blind region for the instrument available at the time of the measurements was around 400 m not allowing to further investigate this point. Over the next day (Fig. 9c), the color-map again indicates the presence of dust from 9:00 to 12:00 at similar heights to 22 June but with lower density until it disappears after 12:00.

720





722

Figure 9: Fine mode fraction as retrieved from CIAO photometer measurements related to the second half of June (a), color-coded time series of range-corrected lidar signals measured at 532 nm cross-polarized channel obtained with the MUSA lidar system on 22 June 2023 (b) and 23 June 2023 (c), aerodynamic particle size distribution daily averages obtained with APS on 22-25 June 2023 (d).

729 Online observations at the ground allow a better understanding of the dust presence at the surface exploring also the status 730 after the 22-23 June. In that period, only APS and aethalometer were operational at CIAO.

Fig. 7d shows the aerodynamic particle size distribution daily averages obtained with APS on 22-25 June 2023 and provides

complementary information to that obtained through lidar and photometer measurements. Indeed, Fig 7d distinctly illustrates

that there is negligible variance in the concentration of ultrafine particulates between dust (22-23 June) and non-dust (24-25

June) days, instead there is a noticeable rise in the concentration of fine and coarse particles with a diameter of up to 5 µm on

the dust days (22-23 June) compared to the non-dust days (24-25 June); demonstrating how during dust events the atmosphere

736	is dominated by large particles (Fig 7a) distributed over different altitude ranges (Fig 7b-c) and if sedimentation is favoured,
737	this leads to a greater return to ground level in the coarse mode (Fig. 7d).
738	This case demonstrates strong agreement between remote sensing and in-situ measurements in identifying and characterizing
739	a desert dust event. Lidar provides detailed vertical and temporal distribution of aerosols, while in-situ measurements capture
740	surface-level dynamics, overcoming lidar's limitations in the blind region near the ground. Together, these methods reveal a
741	more complete picture of the dust event.
742	Future synergistic products could involve integrating additional in-situ instruments with remote sensing to provide richer data
743	for events like this. For example, upgraded systems at CIAO, including advanced offline and online instruments, will enable
744	deeper analyses of similar phenomena. While a detailed investigation of this event is beyond the scope of this paper, it sets the
745	foundation for future studies exploring the full potential of these complementary approaches.
746	More information would be needed for a deeper investigation of such kind of event, and this is the reason why the CIAO
747	observatory has been extensively upgraded as described in this paper and we surely will observe in the next future other events
748	to be analysed through online and offline instruments. However, this kind of detailed investigation is out of the scope of the
749	current paper and would be object of further publications.
750	
751	6 Conclusions
752	The recent upgrade of the CLAO observatory with acrosslip situ laboratory in addition to the well established remote sensing
155	The recent upgrade of the CIAO observatory with aerosof m-situ faboratory in addition to the wen-established remote sensing
754	instrumentation significantly enhances its observational capacity. The integration of in-situ and remote sensing measurements

755 offers a more complete understanding of aerosol behaviour, enabling detailed studies from ground level up to the stratosphere.

756 This combination adds value by providing both vertical profiles by remote sensing measurements and precise ground-level

757 chemical and physical properties through in-situ measurements, which is crucial for improving climate models and

758 <u>understanding aerosol impacts on human health.</u>

Establishing the aerosol in-situ facility has been a complex and labour-intensive endeavour. The process, which began in 2018,
 required careful planning, technical expertise, and collaboration with field specialists. The setup involved designing and
 implementing ACTRIS-compliant inlets, sampling lines, and advanced instruments to ensure accurate and reliable

762 measurements. This development highlights the significant effort required to meet international standards and provide high-

763 <u>quality data for the scientific community.</u>

764 <u>Given CIAO's strategic location in the Mediterranean, the case studies planned for future research are especially relevant. The</u>

765 site is frequently affected by Saharan dust intrusions, which impact air quality and ecosystems, and the observatory is

<sup>766</sup> strategically positioned to study these phenomena. Moreover, the Mediterranean is also prone to wildfires, which are projected

to increase in intensity and frequency due to climate change. The CIAO observatory can monitor both the short-range transport

- of smoke from local fires and long-range plumes from major events, providing insights into their effects on air quality and human health. Lastly, local winter pollution, which results from residential heating, can also be analysed in detail, particularly during temperature inversions that trap pollutants near the ground. The combination of in-situ and remote sensing measurements will help investigate these key environmental issues.
- The recent upgrade of CIAO with the aerosol in-situ laboratory aims to provide comprehensive data on aerosol composition and properties, which will contribute to improve climate change models and understand the effects on human health and ecosystems. The aerosol in-situ laboratory has started in November 2023 the ACTRIS labelling process in order to prove the operational capacities of the National Facility in ACTRIS and ensure the high quality of ACTRIS data in order to obtain the label "ACTRIS National Facility" for the aerosol in-situ component.
- 777 The continuous in situ measurements in tandem with the aerosol remote sensing suite will provide a valuable record of aerosol 778 observations for synergises. Additionally, an ICOS Atmospheric site is under implementation: this will furthermore enhance 779 CIAO's observing capabilities and synergies. All data collected are open and available to external users through international 780 databases (e.g., ACTRIS and ICOS) or through CIAO local services (e.g., meteo data). CIAO also offers remote and physical 781 access to the facility (https://ciao.imaa.cnr.it/access 2/), hosting researchers, students, SMEs and stakeholders, but even the 782 possibility to host user's instruments or sending CIAO mobile platforms to users' sites. All the above are implemented with 783 the main objective of fostering the advancement of the knowledge in the atmospheric field, through the wide use from the 784 scientific community of such extended CIAO observational datasets.
- 785 The CIAO aerosol in situ laboratory has been built following ACTRIS suggestions and requirements, for which technical 786 solutions and schemas are here reported. The instrumental set up will allow to address main research topics such as the aerosol 787 typing and the characterization of the PBL. A first step towards integrating CIAO's different observing platforms is planned 788 during an extensive CIAO measurement campaign focused on the estimation of the PBL using aerosol lidar methodologies 789 and its validation with independent measurements and techniques that will be held in Spring 2024. Furthermore, the next-to-790 come ICOS Atmospheric Class 1 site at CIAO (first step of labelled process already passed) will offer other possibilities of 791 synergistic studies and integration among Ris-RIs in the environmental filed field. In this direction, CIAO is deeply involved 792 in the developments of ITINERIS (Italian Integrated Environmental Research Infrastructures System), an overarching National 793 project for enhancing the interlinkages of all the Italian Ris in the environmental domain. The multi-platform and multi-794 disciplinary approach of the observatory coupled with the open data and open access philosophy is key for better addressing 795 complex atmospheric and environmental questions posed by climate change and anthropization processes.
- 796

797 <u>Appendix A</u>

798Glossary of acronyms

- 799 <u>AAE: Ångström absorption exponent</u>
- 800 ACTRIS: (Aerosol Clouds and Trace gases Research InfraStructure)
- 801 <u>AE: Ångström exponent</u>
- 802 <u>AERONET: Aerosol Robotic NETwork</u>
- 803 <u>APS: Aerodynamic Particle Sizer</u>
- 804 <u>BB: Biomass Burning</u>
- 805 BC: Black Carbon
- 806 <u>CCD: Charge Coupled Plasma</u>
- 807 CIAO: (CNR-IMAA (Consiglio Nazionale delle Ricercahe-Istituto di Metodologie per l'Analisi Ambientale) Atmospheric
- 808 <u>Observatory</u>)
- 809 <u>CNN: Cloud Condensation Nuclei</u>
- 810 <u>CPC: Condensation Particle Counter</u>
- 811 <u>DMA: Differential Mobility Analyzer</u>
- 812 EARLINET: European Aerosol Research Lidar Network to Establish an Aerosol Climatology
- 813 <u>eBC: equivalent Black Carbon</u>
- 814 EC: Elemental Carbon
- 815 <u>FTIR: Fourier Transform Infrared Spetroscopy</u>
- 816 Galion: GAW Aerosol Lidar Observation Network
- 817 GAW: Global Atmosphere Watch
- 818 GCOS: Global Climate Observing System
- 819 <u>GRUAN: GCOS Reference Upper-Ait Network</u>
- 820 <u>HYSPLIT: Hybrid Single-Particle Lagrangian Integrated Trajectory</u>
- 821 ICOS: Integrated Carbon Observing System
- 822 ICP-OES: Inductively Coupled Plasma Emission Spectrophotometer
- 823 <u>NDIR: Non-Dispersive Infrared</u>
- 824 OC: Organic Carbon
- 825 <u>PBL: Planetary Boundary Layer</u>
- 826 <u>PIXE: Particle Induced X-ray Emission</u>
- 827 <u>PLDR: Particle Linear Depolarization Ratio</u>
- 828 <u>PM: Particulate Matter</u>
- 829 <u>RH: Relative Humidity</u>

- 830 <u>RI: Research Infrastructures</u>
- 831 <u>SAE: Ångström scattering exponent</u>
- 832 <u>SMEs: small and medium-sized enterprises</u>
- 833 <u>SMPS: Scanning Mobility Particle Sizer</u>
- 834 <u>TC: Total Carbon</u>
- 835 <u>ToF-ACSM: Time of Flight Aerosol Chemical Speciation Monitor</u>
- 836 WMO: World Meteorological Organization
- 837 <u>XRF: X-ray Fluorescence</u>
- 838

#### 839 Author contributions

840

TL, AM and ST contributed to <u>w</u>Writing – original draft preparation. TL, AM, ST, CCol and MM contributed to <u>v</u>Visualization. TL, AM, ST, FC, DA contributed to <u>m</u>Methodology. TL and MM contributed to <u>f</u>Formal analysis. DA, AG, CD, ER and CC contributed to <u>r</u>Resources. CCor, SG, RMAP, GP contributed to <u>f</u>Funding acquisition. EL contributed to <u>s</u>Software. TL, AM, ST, AA, BDR, MR, LM contributed to <u>w</u>Writing - review & editing. FC, EL, FM, DA, AG, CCor, BDR, CD, SG, MM, NP, GP, RMPA, ER, DS and CCol contributed to <u>r</u>Review & editing. GP and LM contributed to <u>c</u>Conceptualization. LM contributed to <u>p</u>Project administration and <u>s</u>Supervision.

847

#### 848 **Competing interests**

- 849
- The authors declare that they have no conflict of interest.
- 851

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869

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