



Ten years of measurements (2012-2022) of the atmospheric composition at Saclay/SIRTA Observatory in the Ile de France Region as part of ICOS and ACTRIS

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Abstract. CO₂ is the main contributor to global warming, and cities now account for more than two-thirds of emissions of this gas. Atmospheric observatories located on the outskirts of cities are therefore important facilities for measuring the impact on atmospheric composition of the emission reductions planned by cities. The Saclay observatory, part of the ICOS and ACTRIS research infrastructures and located 20 km southwest of Paris, has been monitoring greenhouse gases (CO₂, CH₄), reactive
5 gases (NO_x, O₃, CO), and carbonaceous aerosols (eBC) since 2012. This study presents 10 years of monitoring of these compounds, characterizing diurnal, seasonal cycles and decadal trends. In order to best characterize the impact of Parisian emissions, we defined two sets of data depending on whether the station is downwind of Paris or, conversely, in background conditions with westerly winds. This strategy allows us to characterize the urban offset in the Saclay measurement series. The results show a significant decrease in the urban offset of compounds mainly linked to traffic emissions: -35.6%, -52.3%, and
10 -56.7% for CO, NO_x, and eBC_{lf}. There was also a 15% decrease in urban offset of CO₂ between the 2012-2017 and 2019-2022 periods, a figure consistent with the Airparif inventories' estimate of the decrease in emissions in Paris over the same period.

1 Introduction

Carbon dioxide (CO₂) is a major contributor to global warming which is emitted from various biogenic and anthropogenic
15 sources activities (IPCC, 2022; Friedlingstein et al., 2023). In 2018, almost 55% (4.2 billion) of the population inhabited in urban areas, and according to urban planning projections, in 2050, almost 68% (6.6 billion) of the population will be living within cities (United Nations, 2019). As a result, cities now account for an estimated 70% of global CO₂ emissions and



contribute to anthropogenic CH₄ emissions as well (Mukim, 2023). For this reason, it is becoming increasingly important to quantify those urban emissions, for instance using independent atmospheric observation systems.

20 Long-term measurements of greenhouse gases (GHGs) are relatively rare in urban and peri-urban environments. Atmospheric studies have endeavoured to estimate urban CO₂ emissions. The study by Levin et al. (2011) have been analyzed CO₂ and CH₄ measurements in Heidelberg, Germany over nearly 15 years to estimate emissions with the so-called Radon Tracer Method. They also used ¹⁴CO₂ measurements to separate biogenic and anthropogenic contributions to total CO₂ emissions. In another study by Bezyk et al. (2023), CO₂ and CH₄ concentrations were measured at six stations near the city of Wroclaw
 25 in Poland over a year. That study aimed to align a long-term model with a spatial scale granularity for CO₂ and CH₄. Isotope measurements were also conducted to differentiate between biospheric (emitted by biological processes such as respiration or photosynthesis) and anthropogenic emissions. In a subsequent study by Sreenivas et al. (2016), CO₂ and CH₄ concentrations were measured during one year in India to investigate seasonal impacts. NO_x and O₃ concentrations were also measured to assess the methane sink mechanism. Building upon these earlier findings, Sreenivas et al. (2022) conducted a follow-up study
 30 at the exact location, this time over 3 years. The study's primary objective was to examine and contrast inter-annual variations in CO₂ and CO concentrations with the aid of modelling. Other studies have been conducted using the eddy covariance (EC) method. In Beijing, in the study by Liu et al. (2020), CO₂ and PM_{2.5} were measured between 2009 and 2017 and subsequently studied using the EC method. A study in Sele, Switzerland, was also conducted over 10 years to study trends between 2005 and 2015 (Schmutz et al., 2016).

35 The Paris region is France's most densely populated region, with around 12 million inhabitants, i.e. approximately 20% of the French population over 2% of the country's surface area. Located in a suburban area 20 km southwest from downtown Paris, the Saclay plateau is suitable for monitoring the impact of urban emissions on the regional atmospheric composition, thanks to two observatories associated to the ICOS and ACTRIS European research infrastructures. The ICOS station measures greenhouse gases such as CO₂, CO and CH₄ since 2001. The SIRTa observatory (Site Instrumental de Recherche par Télédé-
 40 tection Atmosphérique, Haeffelin et al. (2005)) linked to ACTRIS measures atmospheric pollutants such as NO_x, eBC and O₃ since 2012/2013. This atmospheric monitoring program, fully involved in the two European Research Infrastructures, provides a unique opportunity to study long-term covariations of greenhouse gases, reactive gases and aerosols in the peri-urban environment of the Paris region. The colocation of these atmospheric measurements is particularly valuable to better characterize emission processes since the observed components are often co-emitted by specific sources like traffic or residential heating.

45 According to the latest inventory, CO₂ emissions in the Paris region were estimated to be 33.5 MtCO₂ in 2021, representing 8.5% of the total emissions in France (Airparif, 2024). The primary sources of CO₂ emissions in the region are transport (31%) and the residential sector (18%) (Airparif, 2024). The main emissions of nitrogen oxides (NO_x) are from the traffic sector (47%), with a smaller contribution from the residential (9%) and tertiary (8%) sectors (Airparif, 2024). The primary sources of carbon monoxide (CO) emissions are the residential sector (55%) and the traffic sector (32%). These two sectors
 50 are also the main sources of black carbon, a particulate pollutant emitted by all types of incomplete combustion (Janssen et al., 2011). Measurement of equivalent black carbon and application of the source apportionment methodology developed by Sandradewi et al. (2008) separate two fractions, the contribution of liquid fuel (eBC_{lf}) and the contribution of solid fuel



(eBC_{sf}), representative in urban areas for the most part of the traffic and residential sectors respectively (Savadkoobi et al., 2023).

55 CO_2 concentration measurements alone are difficult to interpret because the influences of regional emissions are mixed with those on a larger scale, and the influences of biospheric fluxes are mixed with those of anthropogenic emissions from different sectors. The measurement of other compounds used as process tracers, can be used to help discriminate between the different sources of CO_2 variability. To identify the regional contribution to the observed CO_2 variability, the approach is generally to measure atmospheric gradients between two stations located upstream and downstream of the region to determine a baseline from which large-scale contributions can be isolated. In this study, we have developed an alternative method to the gradient approach. We have defined two geographical sectors based on wind direction and speed : one for the urban sector with winds coming from Paris, and one for the rural sector which can be used as a proxy of the background signal. From the difference of the two time series we calculate the urban offset, representing the excess of concentrations that we attribute to the emissions from the Paris area.

65 This study presented atmospheric CO_2 , CH_4 , CO , NO_x , eBC , and O_3 data measurements from 2012 to 2022. In addition to this decade of data, meteorological and boundary layer data and the Airparif inventory for different years were employed. The concentrations are analyzed in the urban and rural sectors (3). These analyses encompassed seasonal (3.1) and diurnal (3.2) variations, culminating in trend analyses across the entire duration of the measurements and the urban offset (3.3). The discussion (4) involves a comparison of the ratios measured with those calculated using the Airparif inventories.

70 2 Materials and Methods

2.1 Site presentation

The Saclay plateau is located in a peri-urban area of the Île-de-France region, 20 km southwest of Paris (France). Measurements of greenhouse gases and pollutants presented in this study come from three locations, 1.5 km apart. CO_2 , CO , and CH_4 were measured at the Laboratory for Climate and Environmental Sciences (LSCE) site from 2012 to 2015 and then at the ICOS tall tower from 2015; meteorological data were also measured on the tower. These measurements are performed within the ICOS European infrastructure and the consistency of the measurements between the two periods has been checked, as presented in the next section (2.2). Measurements of short-lived pollutants (NO_x , O_3 , and BC) were performed at SIRTa in-situ site, located at the LSCE facility. These measurements are part of the ACTRIS (Aerosol, Clouds, and Trace gases Research InfraStructure) European Infrastructure and follow its quality assurance and quality control guidelines (<https://actris.eu/>, last access : 2025/09).

80 Planetary boundary layer height (PBLH) data are issued from measurements at the SIRTa main site, located 5 kilometers from LSCE (Haeffelin et al., 2005; Kotthaus et al., 2023). The measurements between the three sites are consistent because they are located on a plateau and are less than 10 km apart (Table 1). These sites will be referred to as the Saclay site for the remainder of the study.



Table 1. Location and compound used for each site.

Compound	Site	Location
NO _x , O ₃ , BC	SIRTA in-situ	48.709 N, 2.159 E
PBLH	SIRTA main site	48.718 N, 2.208 E
CO ₂ , CO, CH ₄ , meteo	ICOS tower	48.7227 N, 2.142 E

2.2 Instrument presentation

CO₂, CH₄, and CO measurements were made with two gas chromatographs (GC) described by Schmidt et al. (2014) for CO₂ and CH₄ and by Yver et al. (2009) for CO. The GC system was first built in 2001 based on the design of Worthy et al. (1998), and optimized for measurements of CO₂ and CH₄ in flask samples as well as for semi-continuous measurements of ambient air in Saclay. The gas chromatograph used for CO₂ and CH₄ was an HP6890 until 2013 when an HP-7890 replaced it. The GC was equipped with two sample loops and two detectors to analyze simultaneously CH₄ and CO₂ with a flame ionization detector (FID) and N₂O and SF₆ with an electron capture detector (ECD). In July 2006, we coupled a second GC (PP1 analyzer, Peak Laboratories) containing a reduction gas detector (RGD) to the existing GC system to analyze simultaneously CO and H₂. The measurement time step is 5 minutes.

After 2015, CO₂, CH₄ and CO were measured at the ICOS Saclay tall tower by Cavity Ring-down Spectrometers (Picarro G2401 for CO₂, CH₄ and CO, and a Picarro G5310 was installed in 2019 for more accurate CO measurements). Atmospheric air is alternatively sampled and analyzed from the 100m, 60m, and 15m levels in order to get at least one vertical profile per hour, as recommended by the ICOS specifications (ICOS RI, 2020). All of the sampling lines (Synflex1300, 1/2 inch diameters) are connected to pumps in order to ensure a residence time of less than 1 minute. All samples are dried prior to analysis using a Nafion membrane to minimize the water vapor corrections due to spectrometric interferences and atmospheric dilution. The instruments are regularly calibrated against the WMO scales using a set of 4 cylinders bracketing the expected ambient mole fractions. The data quality control is performed by injecting a target gas twice a day to check instrument performance and identify biases.

To correspond as closely as possible to the GC data measured between 2012 and 2015, CRDS data obtained from the ICOS tall tower at three different heights were compared. The data measured at 15 m is used in this article, in combination with GC data sampled from the roof of the LSCE prior to 2015.

As the CO₂, CH₄ and CO data come from several different instruments and locations, we checked the consistency of the dataset. For 2015, the R² results between the GC and Picarro data from the 15 m tower are shown in Figure A1. The R² values are between 0.97 and 0.89 for the three compounds, and the slopes are close to 1. In addition, after 2015, during the lockdown period, the data was measured on the roof of the LSCE by Picarro (Figure A2). For this reason, a correlation was calculated for 2016-2022, with R² values between 0.96 and 0.84 for the three compounds and slopes also close to 1. The results show good correlations between the instruments over the different periods.



NO_x measurements were performed using a chemiluminescence-based instrument equipped with a blue-light converter for converting NO₂ into NO (Teledyne Advanced Pollution Instrumentation (API), model T200UP, USA). Calibration of the instrument is performed monthly, using a 10 ppm primary NO calibration gas standard cylinder (National Physics Laboratory, UK), diluted to 30 ppb with a dynamic dilution calibrator (TELEDYNE API, model T700U) and zero air from a high performance zero air system (TELEDYNE API, model 701H). A gas phase titration (GPT) system, is used every 3 months for the control of the NO₂ converter efficiency. The instrument is equipped with an internal nafion dryer, to avoid the interferences caused by water vapour. The raw signals are corrected for artefacts caused by ozone (when ambient air enters the inlet line, NO₂ photolysis is stopped while the reaction of NO and O₃ continues, resulting in an overestimation of NO₂ and an underestimation of NO). Note that these data are not fully compliant with the ACTRIS measurements guidelines for NO_x, as they have been corrected for ozone but not for the night-time offset. Nevertheless, this correction would be almost always below 0.2 ppb (Robert Wegener, ACTRIS CiGas data QA/QC workshop) and would not affect significantly the results and conclusions presented here.

Ozone measurements were performed with an analyser (Teledyne API, model T400, USA) based on ozone UV absorption (here determined at 254 nm). The instrument has shown a good stability over time and was calibrated at 0 and 100 ppb once a year with a zero-air generator and a calibrator (TELEDYNE API, Model 701H and T700U). The instrument performance has been evaluated in 2017 by an external system from IMT-Douai, giving a value of 0 and 101.1 ppb for zero and calibration respectively. Over the period (2013-2022), the incertitude for O₃ mixing ratio is estimated to be below 10%.

Black Carbon (BC) concentrations were derived from absorption photometer measurements. Single-spot AE31 data were used until February 2013, and compensated using the algorithm presented in Weingartner et al. (2003). Dual-spot AE33 aethalometer has been used since Drinovec et al. (2015). Harmonized quality control has been applied as provided in Petit et al. (2021). Equivalent BC (eBC) mass concentrations were derived following the recommendations of Savadkoobi et al. (2024), by applying a harmonization factor, which depends on aethalometer model and filter tape used, as well as a Mass Absorption Cross-section (MAC) of 7.8 m².g⁻¹ at 880 nm. Source apportioned eBC fractions (liquid fuel and solid fuel) were calculated from the methodology of Sandradewi et al. (2008), with values of α_{sf} and α_{lf} of 1.85 and 0.9, respectively (Petit et al., 2021; Sandradewi et al., 2008).

All times mentioned in this article are expressed in UTC.

2.3 Calculation of the urban offset

The Saclay station is exposed to air masses mainly originating from the South-West sector (the dominant wind as shown in Figure 1), with relatively little urban influence nearby, or to air masses from the North-East that have passed through the highly urbanized Paris region. The hourly measurements in Saclay can thus been classified into two different sectors to analyse the impact of urban contribution on the regional level. The dataset is filtered with meteorological data using wind speed and direction from the top of the Saclay ICOS tower. For the urban sector, the selected wind directions are defined to be between 10° and 70°, corresponding to the north-easterly direction (Figure 1). The rural sector is defined for wind blowing between 190° and 280°, i.e., south-westerly. In order to avoid the impact of nearby cities, a wind speed selection larger than 4 m.s⁻¹ is

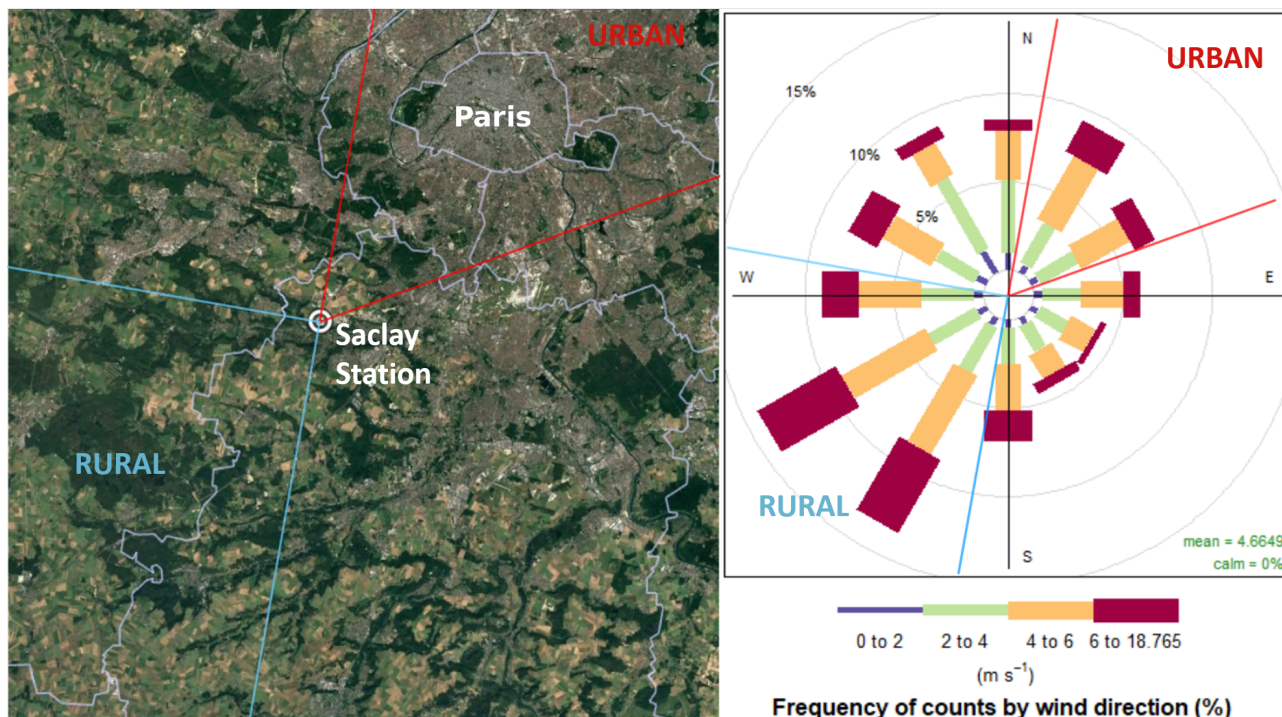


Figure 1. Map of the two geographical sectors used for the classification of the Saclay data. The rural sector is shown in blue and the urban sector in red. The insert shows the wind rose of the dominant wind measured at Saclay. © Google Earth 2019

145 applied on the rural sector. The value of 4 m.s^{-1} is chosen as the best compromise between data availability and the objective of having representative winds. On average, the data from the urban sector represents 15.7% of the total, while the data from the rural sector is 33%. We use measurements from the rural sector to represent background conditions for air masses in the Ile-de-France region. The difference between the two datasets represents the influence of the anthropogenic emissions in the Paris region on the air composition. From the hourly data of the two sectors, we smooth the data using the curve-fitting method of Thoning et al. (1989). This includes a trend function, a series of annual harmonics, and a combination of high-pass and low-pass filters to isolate sub-annual and inter-annual variations, respectively. There is no interpolation of the smooth curves for data gaps of more than thirty days.

2.4 Airparif emission inventory and sources

Airparif, the air quality monitoring structure for the Paris region, is building an emission inventory for the entire Paris region based on a combination of benchmark emission factors and activity data for about 80 emissions sectors. The latest version of the inventory is available for the years 2005, 2010, 2012, 2015, 2019 and 2022 at a $3 \times 3 \text{ km}$ resolution and is based on the updated 2023 methodology. A more detailed description of the Airparif emissions inventory can be found in Bréon et al. (2015). Table 2 shows total CO_2 , CH_4 , CO and NO_x emissions for 2022, as well as the percentages for the main sectors of



Table 2. Contribution of the different emission sectors of compounds in the Ile de FranceParis region according to the Airparif 2022 inventory and total of emissions.

Sectors	CO ₂	CH ₄	NO _x	CO	BC
Residential	20%	18%	8%	65%	70%
Traffic	36%	4%	50%	20%	19%
Waste	12%	55%	3%	1%	0%
Energy	7%	14%	4%	1.4%	0.6%
Tertiaire	13%	1%	8%	1.2%	1.1%
Total	2833 kt	855 t	4193 t	11049 t	143609 kg

activity: residential, traffic, waste and energy. There are two main sources of CO₂ emissions: road traffic, which accounts for a quarter of emissions, and the residential-tertiary sector, which accounts for a third of emissions, mainly from heating. For NO_x, the main emission sector is road traffic, which accounts for almost half of emissions, while for CO, the residential sector accounts for just over half of emissions, and road traffic for a third.

In addition to annual emissions given by sector of activity, Airparif also produces temporal profiles to estimate variations in emissions on a daily, weekly or monthly scale. By combining these temporal profiles, it is possible, for example, to represent typical emissions for winter working days, or summer vacations. Figure 2 shows average diurnal emission profiles for a winter month and a summer month. The road traffic sector unsurprisingly shows strong diurnal variability, for CO₂, CO and NO_x, with very low emissions at night. Emissions rise rapidly between 4 and 8h in response to the onset of urban activity. They then fall slightly, with a small rebound in the late afternoon. Finally, traffic emissions fall towards their residual night values between 17 and 23 h. It is interesting to note that estimated traffic emissions differ very little between summer and winter, both in terms of intensity and diurnal profile. The traffic counting loops to estimate this sector, show that traffic is present and significant throughout the year in the Ile de France region. Conversely, the residential sector, mainly associated with heating, is very low in summer. Winter emissions have daytime profiles similar to those of the traffic sector, with the difference that they extend later into the evening, and only start to fall after 18 h. The double-peak morning/evening emission pattern is clearer in the residential sector than in the traffic sector.

3 Results

Figure 3 shows the complete dataset between 2012 and 2022 with the differentiation between urban and rural sectors. NO, NO₂ and O₃ measurements began in 2013. As expected, concentrations of the compounds mainly emitted by anthropogenic activities (NO_x, BC, CO, CO₂, CH₄) are higher for the urban sector than the rural one due to the influence of the emissions in the Paris region. The variability of those compounds in the urban sector is also more pronounced than for the rural sector. The relatively low concentrations and smaller variability for the rural sector indicates that it represents a good proxy for baseline

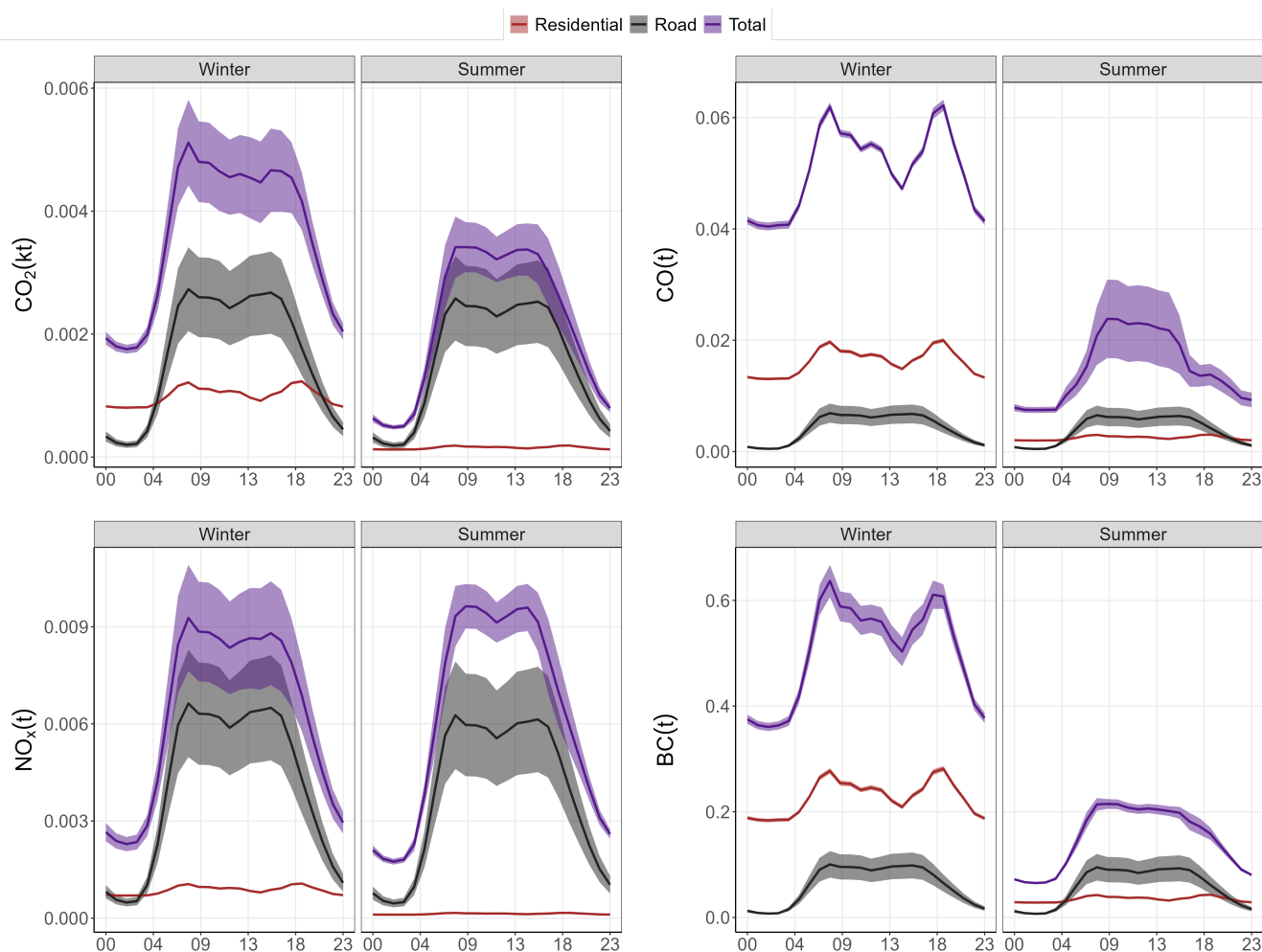


Figure 2. Diurnal cycle in summer and winter using data from the 2022 Airparif inventory for the Paris region for CO_2 , NO_x and CO . The purple curve represents the total inventory, the brown curve the residential and the grey curve the transport. The red curves in the lower panels represent the car count.

conditions in Saclay. We note that for O_3 , a secondary pollutant, the rural sector shows higher concentrations because of the complex mechanics of its formation.

3.1 Seasonal cycles

The mean seasonal cycles for CO_2 , CH_4 , CO , O_3 , NO , NO_2 , eBC_{lf} and eBC_{sf} have been calculated separately for the urban and rural sectors (Figure 4). Concentrations measured in the urban sector are higher than in the rural sector in all seasons and for all compounds, except ozone, which is a secondary pollutant and will be discussed at the end of this section. The



Figure 3. Times Series of eBC_{lf} , eBC_{sf} , NO_2 , NO , O_3 , CO , CH_4 , CO_2 . The black curves represent daily data. The red curve represents the monthly average data for the Urban sector. The blue curve represents the monthly average data for the Rural sector.

differences in concentrations observed between the urban and rural sector depend on emissions and atmospheric dynamics, which change from month to month. The height of the boundary layer is higher in summer, which increases the mixing effect and reduces the measured concentrations at the surface. Conversely, the lower boundary layer concentrates the compounds in winter, increasing the surface concentrations (Foskinis et al., 2024). In addition, emissions from residential heating are higher in winter. According to Kotthaus et al. (2023) boundary layer heights are on average three times higher in summer than in winter at the Saclay site.

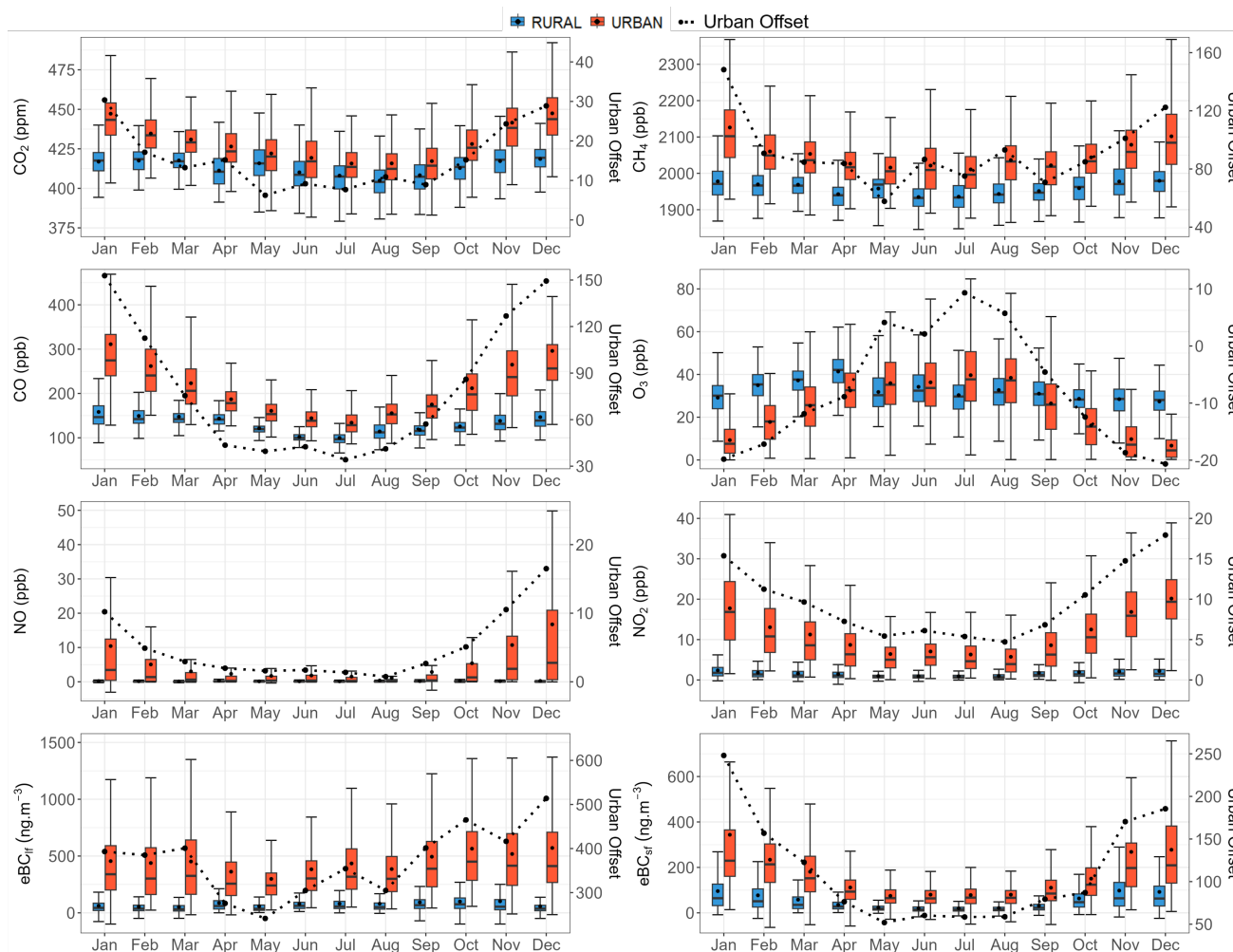


Figure 4. Seasonal cycles have been calculated for all years combined. The red boxplot represents the urban sector, the blue boxplot represents the rural sector. The black curve is the average urban offset represented in the same unit as the corresponding compound. The whiskers extend from the 10th percentile (p10) to 90th percentile (p90), the box from the 25th percentile (p25) to 75th percentile (p75), and the median is plotted by the middle line. The black points represent the mean.

The seasonal cycles of CO₂, CH₄, and CO have similar behaviour in both urban and rural sectors. The seasonal cycles in the two sectors are characterized for all three species by a winter maximum and a summer minimum. These similarities can be explained by common processes associated with anthropogenic emissions and atmospheric transport (Murthy et al., 2020; Foskinis et al., 2024). While atmospheric transport plays an important role in the phase of seasonal gas cycles measured at Saclay, it is not expected to contribute significantly to the seasonal cycle of the urban offset. In fact, the difference in pollutant dispersion between urban and rural areas is not very high, so the seasonal cycle of this signal should more closely reflect that of emissions which are detailed in the following paragraphs.



200 Beyond the seasonal impact of atmospheric dispersion, the seasonal CO₂ cycle is primarily driven by biogenic fluxes, which explains the drop in concentrations between March and August during the plant growth phase, followed by a rise in atmospheric concentrations in autumn and winter when CO₂ respiration by ecosystems exceeds uptake (Watts et al., 2021; Ke et al., 2025). The concentration in the urban sector remains higher due to the influence of the anthropogenic emissions in this sector. The average urban offset is 7 ppm in summer, rising to 25 ppm in winter, giving a ratio of 3.6 between the 2 seasons, 205 which is 20% higher than the ratio given by the Airparif inventory for Paris city.

The average CO urban offset is ranging from 38 ppb in summer to 130 ppb in winter. The winter-to-summer ratio is equal to 3.4, close to the one observed for CO₂, and also about 20% higher than the ratio deduced from the Airparif inventory for Paris city.

The seasonal cycle of atmospheric CH₄ is determined by seasonal variations in sources, chemical loss and transport. In 210 contrast to the national picture, where agriculture is a predominant source, in the Paris region it accounts for less than 10% of CH₄ emissions. Natural sources, mainly associated with wetlands, also play a marginal contribution in this region. CH₄ sinks and atmospheric transport therefore play a dominant role in the seasonal cycle, the main sink being the reaction with the hydroxyl radical (OH), which is produced photochemically and whose local abundance varies seasonally due to the intensity of UV radiation. However, no significant difference in these two processes is expected between urban and rural areas of the 215 Paris region. The seasonal variability of the urban offset, with a ratio of 1.5 between winter and summer, is actually lower for CH₄ than for CO₂ and CO. The same ratio is found in atmospheric observations as in the Airparif inventory, which attributes it to the seasonality of the residential source.

As all reactive compounds, the seasonal cycle of atmospheric NO_x is determined by seasonal variations in sources, photo-chemistry and dynamics. As expected for these very reactive compounds (lifetime of a few hours), which emissions are mainly 220 derived from vehicular traffic, there is a strong difference between levels and variability observed between the urban and the rural sector (Lange et al., 2022). Levels of NO and NO₂ from the rural sector remain below 2.5 ppb the whole year long and show a seasonal variation with a very small amplitude of 0.3 ppb for NO and 1.5 ppb for NO₂. On the opposite, NO and NO₂ from the urban sector (and therefore the urban offset) present a marked seasonal cycle with values 5.2 ppb and 11.2 ppb (on average for NO and NO₂ respectively) higher in winter than in summer. This seasonal cycle can be partly explained by the 225 difference of boundary layer height. Regarding sources, whereas traffic emissions are not associated with significant seasonal variations, residential heating is maximum in winter and contributes to the higher values observed from November to February. The winter/summer atmospheric ratio is 3.7 for NO_x. This is almost 50% higher than the ratio derived from the Airparif inventory for the city of Paris. eBC_{tf} is also mainly emitted by traffic related sources, the seasonality of urban offset is not very pronounced except in autumn following the seasonality of its main source. Thus, it depends strongly on the proximity of the 230 source, which explains the differences in concentrations between urban and rural areas. The urban offset for this compound is less marked than for eBC_{sf}. In contrast, eBC_{sf} is a tracer for the residential heating sector (wood-burning), exhibiting large seasonal variations (Savadkoobi et al., 2024). The increase in concentrations for this compound is evident during the winter period, attributable to the pronounced decline in emissions during the summer months. The urban offset seasonality for eBC_{sf} is pronounced, primarily due to the substantial decrease in emissions in summer.



235 As mentioned above, ozone shows the opposite seasonal pattern to the other compounds, with a maximum in spring/summer and a minimum in winter, in agreement with the seasonal cycle observed in mid-latitudes stations of the northern hemisphere (Schultz et al., 2017). This seasonal variation is mainly driven by the main source of ozone in the lower troposphere, i.e. its photochemical production due to the interplay between precursors (NO_x , CH_4 , CO , VOC) and oxidants (hydroxyl radical). The annual maximum of the urban sector is observed in summer, whereas the one of the rural sector is more shifted towards
240 spring. This last finding is in agreement with the study of Ansari et al. (2024) which have analysed ozone observations over a large number of rural stations in the world. Concerning western Europe and analysing the period (2014-2018) versus (2000-2004), they have noticed a shift of the ozone seasonal maximum towards spring for the most recent period. They explain this summer drop mainly the decrease of NO_x at the national level (due to regulations). Ozone variability associated to the urban sector is more complex to analyse, as it includes local effects (precursors and subsequent photochemistry, ozone titration by
245 NO), see Monks et al. (2015) and references therein. The seasonality of the ozone urban offset is highest in summer (+10 ppb) and lowest in winter (-20 ppb).

3.2 Diurnal cycles

The diurnal cycles of pollutants emitted at the surface are strongly constrained by the dynamics of the atmospheric boundary layer which acts as a key parameter in the dispersion of atmospheric compounds (Foskinis et al., 2024). Consequently, we have
250 analyzed the diurnal cycles of eBC_{lf} , eBC_{sf} , NO_2 , NO , O_3 , CO , CH_4 and CO_2 measured at Saclay, in relation to the boundary layer height measured at the SIRTa main site, 5 km away (Kotthaus et al., 2023). The mean diurnal cycles of all compounds were aggregated by hour, for both rural and urban sectors. For winter, we used the months of December, January, and February, and for summer, the months of June, July, and August (Figure 5).

The first finding on diurnal cycles is that, with the exception of ozone, average concentrations are higher in the urban sector
255 than in the rural sector at all hours of the day. We also note that, in most cases, diurnal cycles are less pronounced in rural than in urban time series, and also in winter compared to summer time.

CO_2 and CH_4 show diurnal cycles with similar behaviors, negatively correlated with PBL heights. Concentrations increase as the boundary layer height decreases at night, and conversely, as the boundary layer expands during the day, CO_2 and CH_4 concentrations decrease. The amplitude of diurnal cycles is more pronounced in summer. It can also be seen that for CH_4 , the
260 summer diurnal cycle is very flat in rural areas, while for CO_2 it is of a similar amplitude to the one in urban areas. This can be explained by the importance of biospheric fluxes in the diurnal CO_2 cycle. The summer maximum of the diurnal cycle for CO_2 and CH_4 in the urban sector is observed between 3 and 5 hr. For most other species (NO , NO_2 , eBC_{lf} , eBC_{sf}), the maximum occurs later, between 5 and 6 hr, and for CO it is even later, between 6 and 9 hr. This period of the day combines the minimum development of the atmospheric boundary layer with the start-up of urban activities and associated emissions, particularly
265 from the urban traffic sector. According to Airparif's emission inventory, CO_2 , CO and NO_x emissions remain high between 6 and 17 hr. So the concentration minima observed in summer are largely explained by the dilution effect associated with the development of the boundary layer. In winter, this dilution effect during the day is much less pronounced and does not generate a morning minimum in atmospheric concentrations, and diurnal cycles therefore have a lower amplitude.

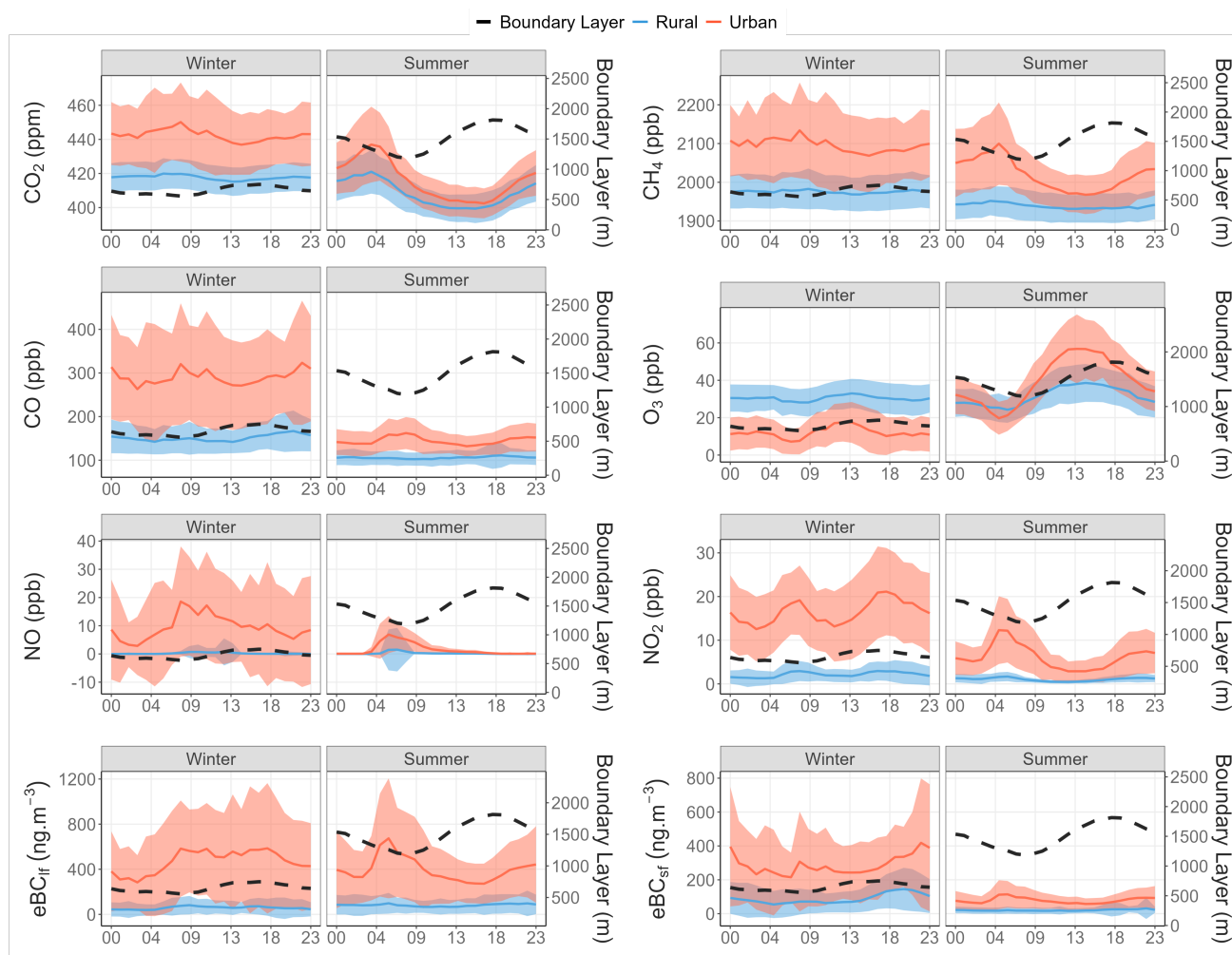


Figure 5. Mean diurnal cycles of concentrations calculated over the period 2012-2022 for Winter and Summer. The red curve represents the average for the Urban sector, with the standard deviation. The blue curve represents the average for the Rural sector. The dotted black curve superimposed on each figure, represents the average boundary layer heights measured between 2015 and 2022 at SIRTa (Kotthaus et al., 2023).

In winter all compounds in the urban sector, except O₃, show a high value peaking at 8 am, due to the high traffic emissions in a low boundary layer. As the boundary layer develops in the course of the day, the concentration of short lived compounds (NO and NO₂) decrease, while concentrations of less reactive compounds remain quite stable. The late afternoon (once the boundary layer height has decreased) is marked by a second peak of NO₂, not observed on NO, this is due to the titration of O₃ by NO. The slightly higher peak of NO₂ in the late afternoon compared to the one observed in the early morning is due to the larger contribution of secondary photochemically formed NO₂, as NO₂ is a primary and secondary pollutant. While traffic is



275 the main source controlling the diurnal cycles of NO_x and eBC_{lf} , the influence of residential heating explains the later evening peak observed on CO and eBC_{sf} .

As mentioned previously, ozone is a secondary pollutant formed by photochemical reactions involving CO, hydrocarbons, NO_x and hydroxyl radicals (Monks et al., 2015). Therefore, its diurnal cycle for the urban sector shows a nighttime minimum and an afternoon maximum; with a larger amplitude in summer, when the photochemistry is maximum. For the rural sector, 280 associated with background concentrations of precursors, the measured ozone hardly shows a maximum during the day, as the observed concentration mainly originates from regional transport.

3.3 Concentration and urban offset trends

As expected, the monitoring of CO_2 and CH_4 concentrations at Saclay reflects the overall long term increase in these greenhouse gases observed at all observatories. For CO_2 , a mean trend of 2.4 ppm per year, during the decade 2012-2022, was 285 observed when using all data, representing an increase of 6% over 10 years. By comparison, the global increase estimated by the National Oceanic and Atmospheric Administration (NOAA) also shows a trend of 2.48 ppm/yr between 2012 and 2022. This was calculated from data available on the NOAA site and using the Mann-Kendall method (<https://gml.noaa.gov/ccgg/trends/index.html>, last access : 2025/09). As CO_2 data have been available since 2003, the slope calculated using the same method is also significant, with an increase of 2.11 ppm per year. By comparison, NOAA data show an increase of 2.27 ppm 290 per year over the same period. For CH_4 we observe a mean annual increase of 10.0 ppb per year, corresponding to a 5% rise over the last decade. This trend is also similar to the global value calculated by the NOAA, which recorded an increase of 9.33 ppb.yr^{-1} between 2012 and 2022. Although we do not observe a significant trend for the last decade for CO, it should be noted that CO data have been available since 2003, and a significant trend of $-4.26 \text{ ppb.yr}^{-1}$ has been calculated when taking into account the whole period (2003-2022). This trend seems consistent with the decrease in anthropogenic emissions over 295 Europe between 2000 and 2017, which is deduced from an inversion of the MOPITT data (Zheng et al., 2019). Concentration of NO_x show a decreasing trend ($-0.17 \text{ ppb.yr}^{-1}$ and $-0.43 \text{ ppb.yr}^{-1}$ for NO and NO_2 respectively) over the studied period (2013-2022). Satellite observations indicate a decrease of NO_2 concentration in cities in developed countries (Erbertseder et al., 2024; Amritha et al., 2024). An overall reduction of NO_x emissions by 18% was also estimated between 2018 and 2023 by a bayesian inversion of TROPOMI NO_2 retrievals (Mols et al., 2025). Over the Paris area a decrease of -2% per year in NO_2 300 levels over the period 1996-2015 was inferred by Erbertseder et al. (2024) and a decrease of 40% of NO_2 has been observed in Paris between 2012 and 2022 (Airparif, 2025). The eBC_{lf} presents as well a negative trend, with a decrease of -24 ng.m^{-3} per year for total data. This downward trend in eBC_{lf} from liquid fuel combustion, mostly traffic, has been observed during the last decade in several European cities (Savadkoohi et al., 2023). With regard to eBC_{sf} from solid fuel combustion, mainly associated with biomass burning, a slight increase in concentrations has been noted since 2020, which could be driven by an 305 increase in wood-burning emissions due to climate policies and rising energy costs (Savadkoohi et al., 2023). The trend calculated for ozone is not significant, in agreement with the observations and simulations reported in the paper by Ansari et al. 2024 for Western Europe. We note that despite a decrease of its precursors (NO_x and VOC), tropospheric ozone is not decreasing.

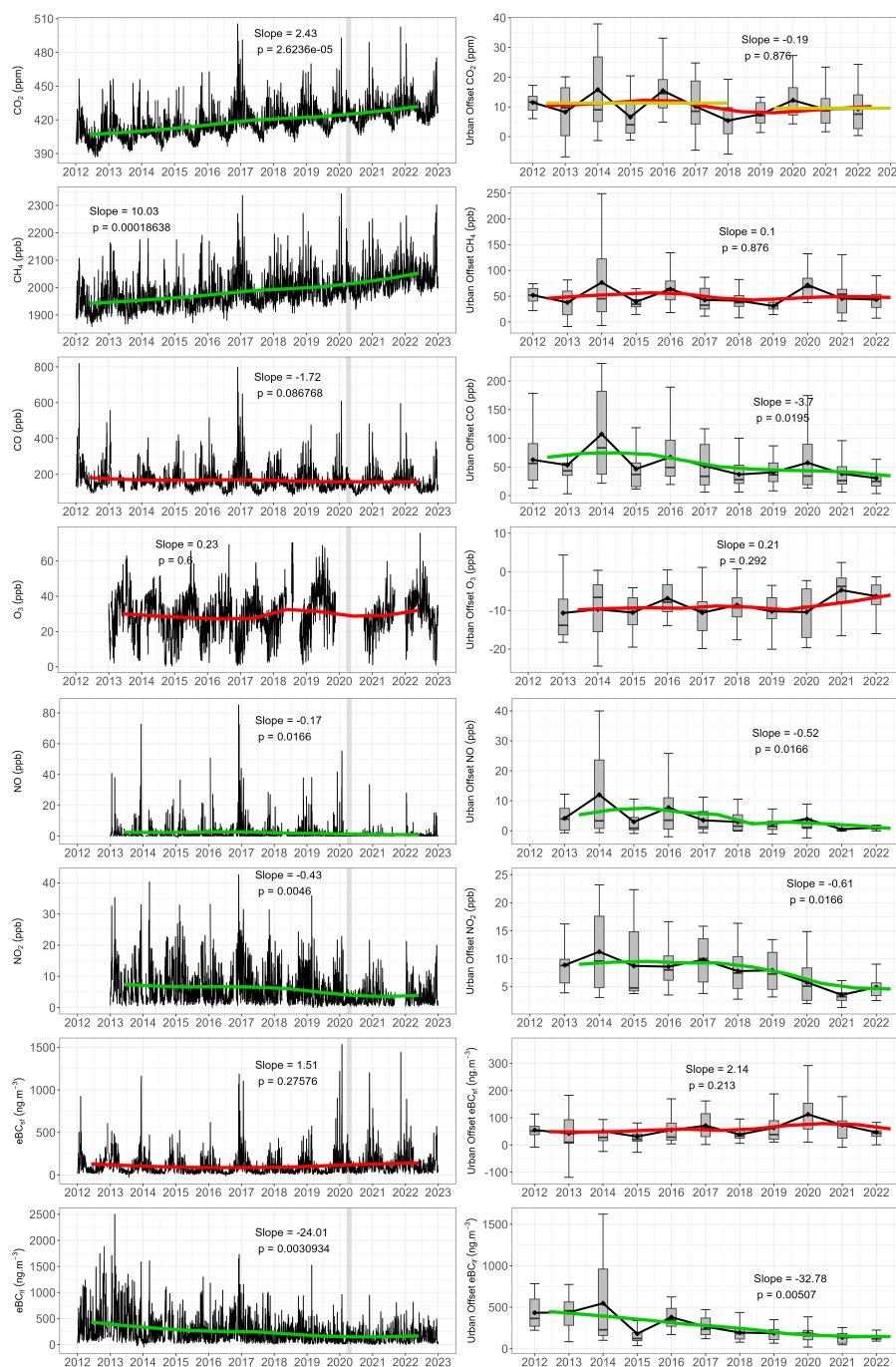


Figure 6. Left panels: daily concentrations (black) and trend calculated using the Mann-Kendall method applied to annual means. Right panel: annual means of the urban offset and trends calculated with the same method. Green curves are significant ($p < 0.05$), and red curves are not significant. The yellow lines for CO_2 urban offset represent the mean values calculated between 2012–2017 and 2018–2022. A vertical grey band identifies the period of the first COVID-19 lockdown in Spring 2020.



Table 3. Summary of the p-values and the slope of the trends between 2012 and 2022 for all the data and the urban offset. Slopes are not considered significant when $p < 0.05$.

Pollutant	All data p-value	All data slope	Urban offset p-value	Urban offset slope
CO ₂	2.62e-5	2.43 ppm·y ⁻¹	0.876	-0.19 ppm·y ⁻¹
CH ₄	1.86e-4	10.0 ppb·y ⁻¹	0.876	0.1 ppb·y ⁻¹
CO	8.67e-2	-1.7 ppb·y ⁻¹	1.95e-2	-3.7 ppb·y ⁻¹
O ₃	0.6	0.2 ppb·y ⁻¹	0.292	0.2 ppb·y ⁻¹
NO	1.66e-2	-0.2 ppb·y ⁻¹	1.66e-2	-0.5 ppb·y ⁻¹
NO ₂	4.60e-3	-0.4 ppb·y ⁻¹	1.66e-2	-0.6 ppb·y ⁻¹
eBC _{lf}	3.09e-3	-24.0 ng·m ⁻³ ·y ⁻¹	5.07e-3	-32.8 ng·m ⁻³ ·y ⁻¹
eBC _{sf}	2.75e-1	1.5 ng·m ⁻³ ·y ⁻¹	2.13e-1	2.1 ng·m ⁻³ ·y ⁻¹

This is due to combined effects of the non-linear chemistry involved in ozone formation, the impact of climate warming and of long-range transport.

310 Concentration trends for long- and medium-lived compounds are representative of a large spatial scale, larger than that of the Paris region. In order to link concentrations time series observed at Saclay to emissions in the Paris region, we also calculated trends in urban offsets. The trend for the urban offset of CO₂ is not significant; however, the average concentration decreases over two periods: between 2012 and 2017, the average concentration of urban offset was 11.3 ppm, while between 2019 and 2022, it was 9.6 ppm, which represents a 15% reduction. These two periods were chosen because there appears to be

315 a break between the average values for these two periods. The urban offset of CO, NO_x, and eBC_{lf} shows significant negative trends of -3.7 ppb.yr⁻¹ for CO, -0.5 ppb.yr⁻¹ for NO, -0.6 ppb.yr⁻¹ for NO₂, and -32.8 ng.m⁻³.yr⁻¹ for eBC_{lf}. All those decreasing trends, deduced from the urban offset, and consequently more representative of the Paris region, are higher than the trends calculated directly with all the concentrations. If we compare the average values of offsets over the periods 2012-2017 and 2019-2022, we have decreases of -35.6%, -52.3% and -56.7% for CO, NO_x and eBC_{lf} respectively. In contrast, eBC_{sf}

320 indicates a 45.7% increase. According to Airparif inventories, the emissions of CO and NO_x have been reduced respectively by -26% and -40% from 2012 to 2021, with a reduction of -50% of the traffic sector for both gases. We do not have the estimates for black carbon. The urban offset calculated from atmospheric measurements in Saclay indicates a higher decrease than the Airparif emission inventory, but with consistency between CO and NO_x. The reduction ratio for these 2 species measured at Saclay is 0.68, close to the ratio of 0.63 for the proportion of traffic in total emissions of these same species (29.4% for CO,

325 46.9% for NO_x).



Figure 7. Ratio of emissions in IDF (diamond) and concentrations (circles) of CO and CO₂ (top), and NO_x and CO₂ (bottom). NO_x and CO (t) and CO₂ (kt).

4 Discussion

Atmospheric CO₂ shares common combustion sources with other compounds like CO, NO_x and eBC, so ratios of measured atmospheric concentrations can help characterize combustion-related CO₂ emissions (Ammoura et al., 2014; Park et al., 2022). In this study, we compared CO/CO₂ and NO_x/CO₂ ratios inferred from atmospheric records to ratios of the same compounds deduced from emission inventories. We have characterized the decadal trends in the ratios between these co-emitted species, as seen from Airparif inventories, and from the measurement of the urban offset of these compounds in Saclay. For the Airparif inventory, we used the 2025 update, which provides emissions estimates for reference years 2012, 2015, 2019, and 2022, based



on the 2023 methodology. We analyse the residential and traffic emission sectors, as well as total emissions of CO_2 , NO_x and CO for the Paris area. As far as atmospheric concentrations are concerned, we take into account the urban offsets calculated as described in section 2.3.

Airparif's inventories, updated every 3 to 5 years, show important decreasing trends of the ratios associated to the traffic sector between 2012 and 2022 (Figure 7). Thus, the ratios of CO/CO_2 and NO_x/CO_2 for traffic-related emissions are declining respectively by 56% and 31%. This reduction is mainly due to the renewal of the vehicle fleet with lower emissions vehicles (following improvements in engine combustion efficiency, in line with Euro standards) and, to a lesser extent, to the reduction in traffic volume following the introduction of some circulation restrictions (Airparif, 2025). For the residential sector, NO_x/CO_2 ratios decrease by 19%, while CO/CO_2 ratios increase by 29% (Airparif, 2021). In terms of total emissions ratios, this translates into a slight drop of 25% for CO/CO_2 and a drop of 31% for NO_x/CO_2 , linked almost exclusively to the traffic sector.

Figure 7 shows that atmospheric ratios of the urban offset are lower than the ratio deduced from the inventory, both for CO/CO_2 and NO_x/CO_2 ratios. One of the reasons can be due to the impact of the biospheric fluxes which are not taken into account in the inventory, but have an impact on the atmospheric CO_2 concentrations. Regarding the decadal trend, the atmospheric ratios also show a decline between 2012 and 2022, of 39% for CO/CO_2 and 47% for NO_x/CO_2 . These decreases are more pronounced than in the inventory, but, like the inventory, show higher values for NO_x/CO_2 than for CO/CO_2 . For NO_x/CO_2 ratios, the annual ratios for the urban offset measured at Saclay are higher than the ratio of emissions from the residential sector, but lower than the ratio of total emissions. The atmospheric CO/CO_2 ratios are close to the total emissions ratio between 2012 and 2018, and after this period, they are very similar to the emissions from the traffic sector.

For the NO_x/CO_2 ratio, one might expect atmospheric observations to correspond more closely to the emissions ratio from the transport sector. However, part of the differences can also be explained by the fact that the measurements are based on a calculation (urban offset) linked to the station's peri-urban location, rather than on direct measurements right at the location of the emission. In particular, determining a delta value for reactive compounds such as NO_x could lead to underestimating the NO_x delta between the emission and the background. Indeed, when comparing the NO_x delta measured at Saclay with that measured at Airparif stations located in Paris or closer to Paris, we observe a higher value for the NO_x delta at these stations. This suggests that such comparisons should favor long-lived compounds such as CO or stations closer to the urban center.

5 Conclusions

This study analyzes ten years of atmospheric CO_2 concentrations, co-located with measurements of various gases and aerosols (CH_4 , CO, O_3 , NO_x , eBC_{lf} and eBC_{sf}) at the peri-urban site of SIRTSA/Saclay, in the Paris region. Due to the location of the measurement station, 20 km South-West of Paris, we distinguish rural (representative of background air) and urban sectors and calculated the corresponding urban offset in the atmospheric time series observed at Saclay.

The observed time series were interpreted using diurnal cycles by season, as well as seasonal cycles. Both showed a substantial impact of biogenic emissions on CO_2 . Consequently, the CO_2 data appear to be less correlated with the other compounds, mostly emitted by anthropogenic sources, during the summer when biogenic fluxes are at their maximum. In addition, the



height of the boundary layer during this period also significantly impacts the concentrations measured for atmospheric pollutants, with the concentrations measured during this period being lower than in winter. The decadal trends for CO₂ and CH₄ concentrations show similar increases to other monitoring stations worldwide over the same period. In contrast, some compounds (NO_x; eBC_{lf}) show decreasing trends when all the data are considered. The analysis of the urban offset allows us
 370 to focus more on the impact of activities from the Paris region, since we subtract a background signal, which is particularly required for species with a long lifetime. We found significant downward trends for compounds related to emissions from the traffic sector, namely CO, NO_x and eBC_{lf}. If we compare the urban offset from the 2012-2017 period to the 2019-2022 period, we observe decreases of -35.6%, -52.3% and -56.7% respectively, for CO, NO_x and eBC_{lf}. For its part, CO₂ offset shows a 15%, comparable to the decrease in anthropogenic CO₂ emissions in the Paris region according to Airparif inventories for the
 375 same period.

A comparison of NO_x/CO₂ and NO_x/CO₂ ratios with Airparif inventories reveals similar downward trends, more pronounced in atmospheric observations. The atmospheric ratios of CO/CO₂ and NO_x/CO₂ deduced from the urban offset are lower than the ratios deduced from the inventory. This underestimation can probably be partially explained by the role of biospheric fluxes that are not taken into account in the inventory. The NO_x/CO₂ ratio from the Saclay observatory may also be
 380 underestimated, as NO_x are reactive species that rapidly transform in the atmosphere. Moreover, it is important to note that the species ratios are not constant over time; they evolve with changes in technology, regulations, and fuel composition. Therefore, using NO_x or CO as anthropogenic combustion tracers for atmospheric CO₂, requires accounting for these long-term trends to ensure an accurate interpretation.

The long-term monitoring at SIRTAS/Saclay observatory is performed within the European Research Infrastructures ICOS
 385 and ACTRIS framework. It shows the added value of having co-located measurements of greenhouse gases and pollutants, as they share some familiar sources that help to confirm, for example, the role of the traffic sector in the decrease of CO₂ offset.

Data availability. All data used in this article, except for boundary layer height data, are available in the Easy Data catalogue at <https://doi.org/10.57932/5c399263-a317-41b7-8900-184b177c4216> (Bouillon and Ramonet, 2025).

Appendix A

390 A1

Author contributions. methodology, V.G. and M.R.; validation, V.G. and M.R.; formal analysis, L.B.; investigation, L.B.; data curation, C.P., C.Y.K., L.D. O.P., J-E.P. and N.B.; writing—original draft preparation, L.B., M.R. and V.G.; writing—review and editing, all; supervision, V.G. and M.R.

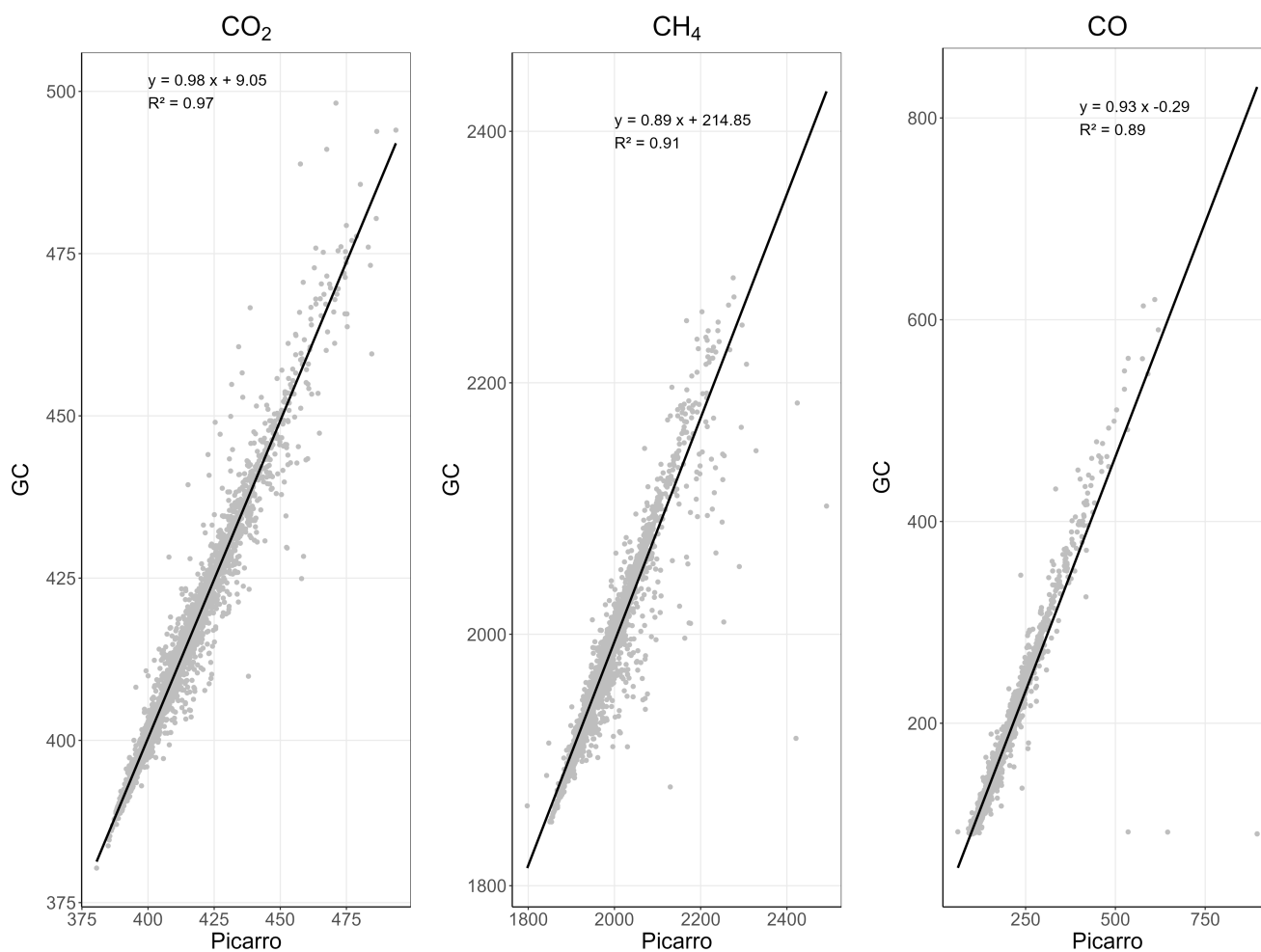


Figure A1. Correlation between data measured by the GC located at LSCE and data measured by the Picarro at the Saclay tower at 15m between 2015-01-01 and 2015-12-31

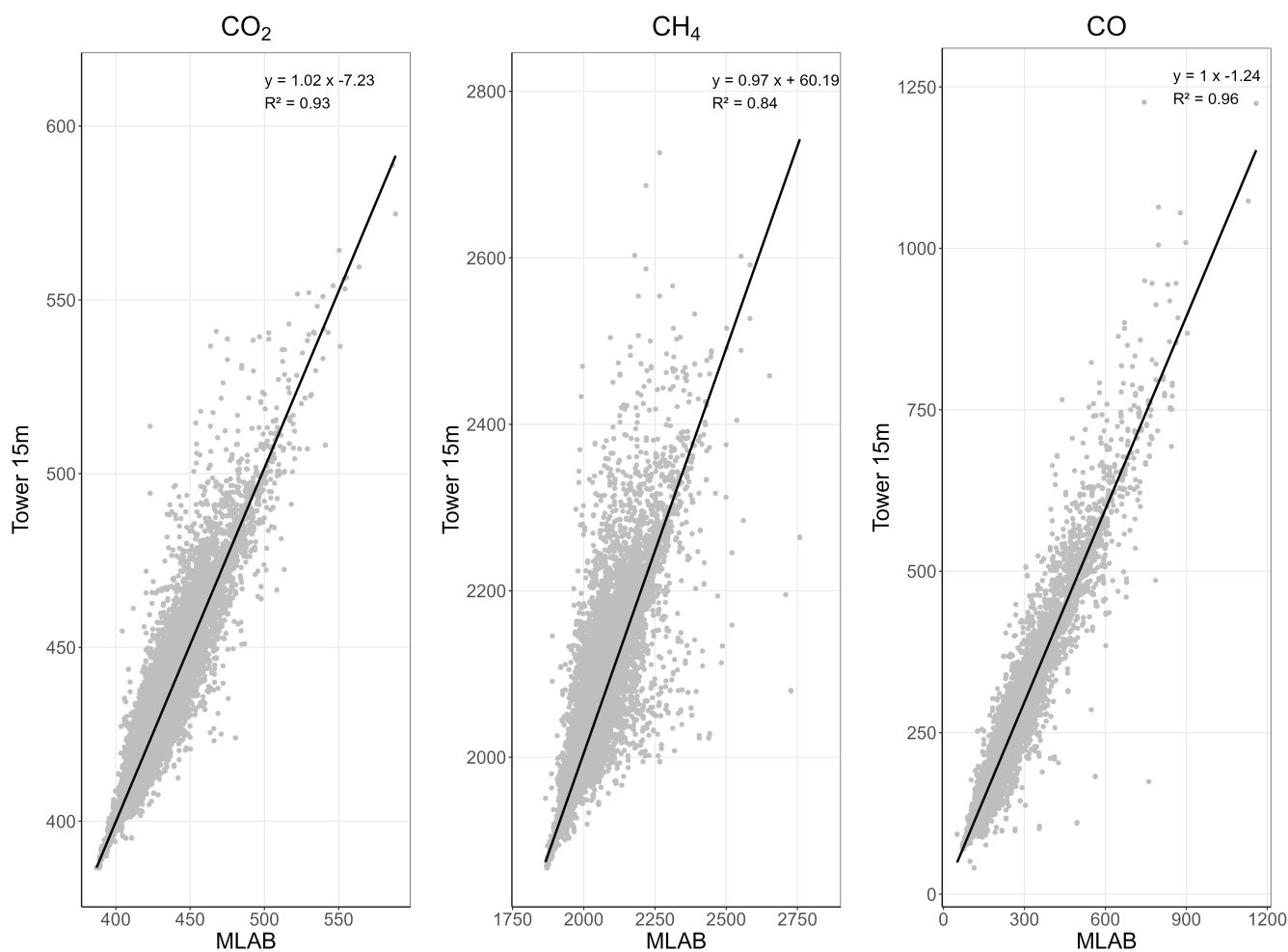


Figure A2. Correlation between data measured by the Picarro located at LSCE and data measured by the Picarro at the Saclay tower at 15m between 2016-01-01 and 2022-12-31



Competing interests. The authors declare that they have no conflict of interest.

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