Analyzing Four Years of Ground-Based Measurements of XCO₂ and XCO over Thessaloniki, Greece Using FTIR Spectroscopy †

Thomas Panou 1,*, Chrysanthi Topaloglou 1, Marios Mermigkas 1○, Dimitrios Balis 1○, Darko Dubravica 2 and Frank Hase 2

1 Laboratory of Atmospheric Physics, Department of Physics, Aristotle University of Thessaloniki, Campus Box 149, 54124 Thessaloniki, Greece; chtopal@auth.gr (C.T.); mmermigk@physics.auth.gr (M.M.); balis@auth.gr (D.B.)
2 Institute of Meteorology and Climate Research (IMK-ASF), Karlsruhe Institute of Technology, P.O. Box 3640, D-76021 Karlsruhe, Germany; darko.dubravica@kit.edu (D.D.); frank.hase@kit.edu (F.H.)
* Correspondence: tpanou@physics.auth.gr

Abstract: The issue of atmospheric pollution in urban centers has become a growing concern in recent years. The increasing levels of greenhouse gases in the atmosphere are a major contributor to atmospheric pollution, and it is imperative to monitor these gases. This study presents the measurements of column-averaged dry-air mole fractions of carbon dioxide (XCO₂) and carbon monoxide (XCO) in Thessaloniki, Greece. The measurements were taken in Thessaloniki using the Bruker EM27/SUN instrument, which was developed by Bruker and KIT and has been part of the Collaborative Carbon Column Observing Network (COCCON) since 2018. COCCON is a global network of stations around the globe and serves as an important supplement to the high-resolution Bruker IFS125 spectrometer used in the Total Carbon Column Observing Network (TCCON), and it provides an increased density of column-averaged greenhouse gas observations. In this work, a four-year analysis of column-averaged dry-air mole fractions of XCO₂ and XCO is presented, focusing on diurnal and seasonal cycles as well as on the comparison between them. The hourly time series show the expected seasonal cycle of XCO₂ with a spring maximum and late summer minimum due to photosynthesis activity, while XCO₂ presents a daily maximum of 419.987 ± 2.286 ppm and a daily minimum of 405.001 ± 3.067 ppm. The seasonal co-variability between XCO₂ and XCO reveals an interesting correlation—especially during winter (R² = 0.841 for 2022) and spring (R² = 0.437 for 2022) period, when anthropogenic emission sources occur.

Keywords: FTIR; EM27/SUN; COCCON; Thessaloniki; XCO₂; XCO; KIT; residuals; seasonality

1. Introduction

Climate change is one of the most pressing challenges our society is facing today and for generations to come. The use of fossil fuels is the primary anthropogenic source contributing to the carbon cycle, along with the development of the agricultural industry, coal mining, waste management, and other human activities that lead to increased concentrations of greenhouse gases [1,2]. The resulting rise in greenhouse gases, particularly carbon dioxide (CO₂), is driving climate change, including the greenhouse effect and global warming.

CO₂ is a well-mixed gas that has a longer lifetime and contributes approximately 65% of the radiative forcing responsible for climate change. Despite global efforts to reduce emissions, its concentrations are increasing at the fastest observed decadal rate of change (2.0 ± 0.1 ppm/yr) for the period of 2002–2011 [3]. Carbon monoxide (CO) is primarily sourced from the oxidation of methane, biomass burning, and fossil fuel combustion. The primary sink of CO in the atmosphere is through oxidation with hydroxyl radicals.
(OH) [4]. Therefore, it is essential to measure greenhouse gas concentrations and study their spatial and temporal variations accurately and precisely. This information is crucial for estimating global emission levels and improving our understanding of their sources and sinks, ultimately informing policy decisions aimed at mitigating global warming.

Various methods can be employed for the measurement of greenhouse gases, including in situ measurements that offer insights into local sources and satellite remote sensing. In this study, we utilize Fourier Transform Infrared Spectroscopy (FTIR) to determine the total column abundance of greenhouse gases across all atmospheric layers. On a global scale, in situ measurements are carried out using FTIR spectrometers that use direct solar radiation as a light source. The Total Carbon Column Observation Network (TCCON) [5], established in 2004, is designed to obtain accurate and precise column abundances of greenhouse gases, such as CO₂, CH₄, and CO, from near-infrared solar absorption spectra. TCCON stations operate high-resolution Fourier Transform IFS125HR spectrometers (FTS) manufactured by Bruker [5]. In 2016, the COCCON network emerged as a viable alternative, based on the low-resolution (0.5 cm⁻¹) EM27/SUN FTIR spectrometer developed by KIT in collaboration with Bruker [6].

2. Methodology

The initial data were divided into hourly, daily, and monthly averages for analysis. The primary objective was to determine the co-variability of XCO₂ and XCO gases and to identify their common sources. These gases displayed irregular seasonal fluctuations, so hourly mean concentrations per year were calculated and subtracted from the original data to remove seasonal variations. This filtering process enabled the extraction of short-term variations caused by local effects for each gas, referred to as ΔX terms or residuals. The resulting data present short-term fluctuations that can be examined to better understand each gas’s behavior.

\[ \Delta X_{\text{gas}} \text{ or Residual}_{\text{gas}} = X_{\text{gas}} - X_{\text{gas \ annual \ cycle}} \]  

After identifying the X gases for each season, we conducted a comparison of residuals to investigate their seasonal and daily relationships.

3. Results

In this section, the described methodology is applied using FTIR spectrometer observations with the aim of extracting the residual values of XCO₂ and XCO, in order to establish a connection between their common sources and sinks. The residual values correspond to the differences between the observed concentrations and the concentrations that would be expected based on known sources and sinks. By identifying and analyzing these residual values, we can gain insights into the underlying processes that contribute to changes in atmospheric XCO₂ and XCO. This analysis may have important implications for understanding the dynamics of the Earth’s carbon cycle and for predicting future changes in atmospheric composition.

Carbon Monoxide and Carbon Dioxide Co-Variability

Table 1 presents the average annual values for the four-year time series. The annual increase in carbon dioxide is evident mainly due to anthropogenic emissions, in contrast to XCO, which shows a comparatively more stable trend with occasional increases caused by pollution events such as fires.

<table>
<thead>
<tr>
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<th>2019</th>
<th>2020</th>
<th>2021</th>
<th>2022</th>
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<tr>
<td>XCO₂ (ppm)</td>
<td>409.59 ± 1.93</td>
<td>412.41 ± 2.29</td>
<td>414.65 ± 2.53</td>
<td>416.97 ± 1.73</td>
</tr>
<tr>
<td>XCO (ppm)</td>
<td>0.088 ± 0.007</td>
<td>0.091 ± 0.008</td>
<td>0.097 ± 0.009</td>
<td>0.092 ± 0.008</td>
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Figure 1a presents the daily and monthly average time series of XCO₂ for the four years. The data reveal a clear seasonal cycle in the XCO₂ concentrations, which is primarily caused by the natural photosynthesis cycle. During the Northern Hemisphere spring and summer, plants absorb CO₂ from the atmosphere for photosynthesis, leading to a decrease in atmospheric CO₂ concentrations. Conversely, during fall and winter, plants shed their leaves and release CO₂ back into the atmosphere, causing an increase in atmospheric CO₂ concentrations.

![Figure 1a](image1.png) ![Figure 1b](image2.png)

**Figure 1.** (a) XCO₂ daily mean of the dry-air column with its corresponding temporal standard deviation, with the trend of monthly mean. (b) XCO daily mean of the dry-air column with its corresponding temporal standard deviation, with the trend of monthly mean.

In addition to the natural seasonal cycle, Table 1 shows that there has been an overall increase in atmospheric CO₂ concentrations over the four-year period. This increase is mainly attributed to human activities, particularly the burning of fossil fuels and deforestation. These activities release large amounts of CO₂ into the atmosphere, which accumulate over time and contribute to global warming and climate change.

The time series data for carbon monoxide (CO) presented in Table 1 and Figure 1b show a relatively stable picture over time, with CO concentrations remaining relatively consistent throughout the four-year period. The concentration of carbon monoxide often displays several gaps, particularly in the summer months, as a result of pollution episodes. These episodes are primarily caused by fires, such as the ones that occurred in July 2021.

The correlation between ΔXCO₂ and ΔXCO values in Figure 2 suggests that changes in fossil fuel combustion can affect the levels of both gases. It is important to note that other factors, such as natural emissions and atmospheric processes, can also contribute to variations in ΔXCO₂ and ΔXCO values. However, the strong correlation, especially in the winter months, between these two gases indicates that they have common sources.

![Figure 2](image3.png)

**Figure 2.** (a) ΔXCO values in Figure 2 suggests that changes in fossil fuel combustion can affect the levels of both gases. (b) ΔXCO values in Figure 2 suggests that changes in fossil fuel combustion can affect the levels of both gases.
The time series for $XCO_2$ demonstrates the emergence of the seasonal cycle, with maxima occurring in winter ($423.093 \pm 2.817$ ppm) and spring ($417.031 \pm 2.863$ ppm) due to anthropogenic influence and a summer minimum ($412.316 \pm 3.067$ ppm) resulting from the biosphere cycle, which is a local phenomenon for mid-latitude regions of the Northern Hemisphere. The time series for $XCO$ exhibits a greater variability in its daily values due to local sources, with a maximum ($0.120 \pm 0.006$ ppm) occurring in winter and a minimum in summer ($0.070 \pm 0.009$ ppm), reflecting the common sources shared with carbon dioxide. However, the concentration of carbon monoxide can sometimes reach high levels during the summer months due to fires occurring in those periods, as was the case in July 2021 when the concentration was measured at $0.141 \pm 0.013$ ppm.

The study of the co-variability of $\Delta XCO$ and $\Delta XCO_2$ residuals throughout the seasonal cycle showed good agreement for the winter months ($R^2 = 0.665$) and a significant correlation coefficient in spring. These findings suggest that there are strong influences of fossil fuel as a common emission source for the two gases during these seasons. However, in the summer and autumn months, the two gases are not correlated due to the higher seasonality of $XCO_2$ compared to $XCO$, indicating that the emission sources for these two gases differ during these seasons. Therefore, it is crucial to consider the seasonal variability of the sources when studying atmospheric chemistry and air pollution.

**Figure 2.** Correlation between collocated $\Delta XCO$ and $\Delta XCO_2$ for winters 2019 (a), 2020 (b), 2021 (c), and 2022 (d), along with the resulting linear correlation line. A faded light-blue color represents the confidence interval.

4. Conclusions

In this study, the total columns of carbon dioxide and carbon monoxide in Thessaloniki, Greece were measured for four full years (2019–2022), using a portable direct solar viewing FTIR spectrometer. Additionally, the seasonal behavior of greenhouse gas residuals ($\Delta X$) was examined to capture short-term variations of X-gases and identify local sources, as well as common sources between them.

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**Data Availability Statement:** The EM27/SUN FTIR spectrometer data over Thessaloniki, Greece are available upon request.
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Conflicts of Interest: The authors declare no conflict of interest.

References

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