

1 Atmospheric CO₂ dynamics in a coastal megacity: spatiotemporal 2 patterns, sea–land breeze impacts, and anthropogenic–biogenic 3 emission partitioning

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15 **Abstract.** Attributing observed carbon dioxide (CO₂) to fossil-fuel emissions versus biogenic fluxes is essential
16 for assessing urban mitigation, but in coastal megacities it is complicated by anthropogenic–biogenic coupling and
17 sea–land breeze (SLB) circulation. Here we analyze Guangzhou using multi-site in situ CO₂ and CO measurements
18 (January 2023–September 2024), transport footprints, and a site-specific $\Delta\text{CO}/\Delta\text{CO}_2$ (R_{CO}) relationship to resolve
19 spatiotemporal variability, quantify SLB effects, and partition fossil-fuel (CO₂ff) and biogenic (CO₂bio)
20 contributions without assimilating emission inventories. Along a coastal–urban–suburban gradient, the coastal site
21 shows the largest seasonal amplitude (25.63 ppm), the vegetated site exhibits strong summertime diurnal amplitude
22 (39.90 ppm), and the urban core is combustion-dominated; together, these gradients indicate a seasonally displaced
23 coastal CO₂ “dome”, whose maximum enhancement need not occur in the urban core. SLB effects are seasonal:
24 SLB ventilates CO₂ in spring–winter but promotes summertime accumulation (+2.08 ppm) under stable
25 stratification, accompanied by pronounced CO enhancements, consistent with trapped/recirculated combustion
26 plumes. Regression-derived urban R_{CO} is consistent with reported post-2013 broad tightening of coal/industrial and
27 vehicle-emission controls. Winter-afternoon urban CO₂ff attribution remains robust to transport-model
28 configurations and quantified measurement/background uncertainty. Summer-afternoon CO₂bio is -3.59 ± 0.63
29 $\mu\text{mol m}^{-2} \text{s}^{-1}$, offsetting up to 60 % of concurrent CO₂ff. These results demonstrate that coastal dynamics and urban
30 greening reshape observed CO₂ signals, highlighting that biogenic–anthropogenic decoupling and SLB-aware
31 sampling are essential for the robust evaluation of carbon mitigation in coastal megacities.

32 1 Introduction

33 Atmospheric carbon dioxide (CO₂), the predominant anthropogenic driver of climate change, is accumulating at
34 unprecedented rates in human history (WMO, 2024). Future CO₂ increments will exert stronger warming effects
35 than equivalent past increases due to climate system feedbacks (He et al., 2023), making emission control
36 imperative. Despite covering only 3 % of global land, urban areas generate over 70 % of carbon emissions (Crippa
37 et al., 2021), positioning cities as critical arenas where mitigation policies, infrastructure transitions, and urban
38 greening can drive measurable changes in urban carbon budgets (WMO, 2025).

39

40 Urban atmospheric CO₂ concentrations provide a complementary constraint to flux- or inventory-based
41 assessments because they integrate surface emissions, biogenic exchange, and atmospheric transport over the
42 upwind source area (Lin et al., 2003; Shusterman et al., 2018; Pitt et al., 2022). This integration, however, creates
43 an attribution challenge because observed CO₂ variability reflects coupled effects of fossil-fuel emissions, biogenic
44 fluxes, and meteorology/transport, and thus is not uniquely attributable to any single driver, although one factor
45 may dominate in certain regimes, especially on diurnal to seasonal timescales (Xueref-Remy et al., 2018; Yang et
46 al., 2021; Mitchell et al., 2018). The challenge is particularly acute in coastal megacities, where land–sea thermal
47 contrasts and marine background inflow drive strong diurnal reversals in advection and boundary-layer structure
48 (Leroyer et al., 2014; Lei et al., 2024). These dynamics complicate observational representativeness and source–
49 sink separation, and can make inferred CO₂ enhancements sensitive to background selection under different
50 transport regimes (Verhulst et al., 2017). Given their high exposure to climate-related hazards, especially flooding
51 and storm impacts, coastal megacities are priority targets for climate-risk management, motivating robust, policy-
52 relevant CO₂ emissions assessment (Kumar, 2021).

53

54 A range of approaches has been used to constrain urban carbon budgets, including bottom-up inventories, tower-
55 based atmospheric observations, tracer/isotope measurements, and atmospheric inversions. Bottom-up inventories
56 provide essential baselines, yet urban emissions remain uncertain and can differ across products because they
57 depend on activity/emission-factor assumptions and proxy-based downscaling (Gately and Hutyra, 2017; Gurney
58 et al., 2019). While radiocarbon (¹⁴C) measurements provide robust fossil-fuel/biogenic partitioning (Turnbull et
59 al., 2015; Berhanu et al., 2017; Wang et al., 2022), their high cost and discontinuous sampling limit their
60 applicability for long-term, high-frequency monitoring. Similarly, eddy covariance (EC) measurements quantify
61 net CO₂ fluxes within tower footprints (typically 1–2 km radius), yet the derived flux partitioning reflects only
62 local-scale dynamics and cannot fully represent city-wide carbon exchange processes (Velasco et al., 2013; Menzer

63 and Mcfadden, 2017; Wu et al., 2022b). Fully three-dimensional (3-D) atmospheric inversions can assimilate
64 concentration data to estimate city-scale emissions and quantify posterior uncertainty when observation networks
65 adequately constrain boundary inflow and key transport uncertainties (Lauvaux et al., 2016; Lian et al., 2023). Yet
66 in practice, urban inversions are often limited by incomplete observational coverage, representativeness (sub-grid)
67 errors, and transport biases (Boon et al., 2016; Deng et al., 2017; Ye et al., 2020; Che et al., 2022b). They can also
68 remain sensitive to choices of background/boundary conditions and prior assumptions, including those related to
69 biogenic fluxes, which can be large and temporally variable during the growing season (Turnbull et al., 2018; Nalini
70 et al., 2022; Ye et al., 2020). These constraints motivate complementary observation-driven frameworks that
71 emphasize process-level interpretation of concentration variability while explicitly tracking uncertainty sources.

72

73 A defining coastal process is the sea–land breeze (SLB), a common mesoscale circulation driven by land–sea
74 thermal contrast (Shen et al., 2021). In the Pearl River Estuary (PRE), SLB occurrence shows pronounced
75 seasonality and spatial dependence, with spring-autumn maxima reported in parts of the region (You and Chi-Hung
76 Fung, 2019; Mai et al., 2024b). SLB produces a clear diurnal reversal between onshore and offshore flow and can
77 modify boundary-layer structure and mixing depth (Wu et al., 2013; Lei et al., 2024), complicating the
78 interpretation of urban CO₂ variability. By contrast, inland basins/valleys lack marine inflow–outflow and may be
79 shaped by lake–land breeze circulations (Yang et al., 2022) or other topography-modulated flows that can shift
80 background conditions and the footprint representativeness of urban observations (Mitchell et al., 2018; Wang et
81 al., 2021). Beyond the PRE, China’s coastal basins exhibit distinct SLB seasonality (e.g., a summer maximum in
82 the Bohai Rim, weak seasonality in the Yangtze River Delta, and an autumn maximum in the PRE) (Huang et al.,
83 2025), likely driven by regional differences in large-scale wind fields and air–sea turbulent heat fluxes (Shen et al.,
84 2024), and potentially modulated by coastline geometry and topographic constraints. Despite recognized impacts
85 of SLB on air quality (Zhao et al., 2022; Wang et al., 2023; Zheng et al., 2024), only a few case studies have
86 examined how SLB-driven recirculation of nocturnally respired CO₂ can perturb daytime coastal CO₂
87 measurements and may bias emission inversions (Ahmadov et al., 2007), and systematic quantification across
88 coastal cities remains limited.

89

90 Urban greening reinforces the need for explicit source–sink separation because neglecting biogenic contributions
91 can bias inferred urban emissions (Miller et al., 2020; Ye et al., 2020; Sargent et al., 2018). Moreover, urban
92 vegetation is often highly fragmented and can be underrepresented in moderate-to-coarse remote-sensing land-
93 cover/canopy products, which may miss fine-scale urban heterogeneity and thereby introduce additional

94 uncertainty (and potential bias) into satellite-driven biogenic flux estimates (Ma et al., 2022; Corro et al., 2025;
95 Glauch et al., 2025). Green coverage within built-up urban areas is similarly high across major Chinese coastal
96 megacities (e.g., Guangzhou: 43.7 % in 2023; Shenzhen: 44.0 % in 2023; Qingdao: 44.0 % in 2023; Tianjin: 38.2 %
97 in 2023) (Guangzhou Municipal Bureau of Statistics, 2024; Shenzhen Municipal Bureau of Statistics, 2024;
98 Qingdao Municipal Bureau of Statistics, 2024; Tianjin Municipal Bureau of Statistics, 2024), implying that
99 biogenic exchange can be non-negligible when interpreting urban observations. For example, in coastal North
100 American cities, high-resolution vegetation modeling in New York City indicates that summertime biogenic uptake
101 can offset up to 40 % of afternoon anthropogenic CO₂ enhancements and can fully balance on-road traffic
102 contributions (Wei et al., 2022), while a dense-sensor analysis in Los Angeles suggests that daytime biogenic
103 exchange can consume up to 60 % of fossil-fuel emissions during the peak growing season (Kim et al., 2025).
104 Recent advances, including emerging atmospheric constraints such as carbonyl sulfide (COS), further demonstrate
105 the feasibility and value of more robust biogenic quantification in city carbon budgets (Soininen et al., 2025).

106

107 Taken together, three knowledge gaps limit the interpretation of coastal megacity CO₂ observations and their use
108 for mitigation assessment: (i) how spatiotemporal CO₂ patterns vary across a coastal–urban–suburban gradient; (ii)
109 how SLB-driven transport and boundary-layer mixing reshape diurnal and seasonal CO₂ signals through seasonally
110 varying ventilation, boundary-layer depth, and atmospheric stability, which may differ from inland-city regimes;
111 and (iii) how fossil-fuel and biogenic contributions can be robustly separated, with quantified uncertainty, in a
112 coastal setting. To bridge these gaps, we investigate Guangzhou (Pearl River Delta, China) as a living laboratory:
113 high GDP and population indicate strong energy and mobility demand and thus substantial fossil-fuel CO₂
114 emissions, while evergreen urban vegetation, a long growing season, and frequent SLB circulation can produce
115 non-negligible biogenic signals and complex coastal transport. The policy context further elevates the value of
116 attribution: Guangdong’s emissions-trading system (ETS) has operated since 2013, alongside measures targeting
117 energy structure, vehicle emissions, building efficiency, and urban greening.

118

119 Given this imperative to track changes in urban emissions and net carbon exchange, we use an observation-driven
120 framework that complements atmospheric inversion approaches and interprets diurnal–seasonal variability by
121 combining multi-site in-situ CO₂ and CO measurements with footprint-informed transport analysis (Lin et al., 2003;
122 Fasoli et al., 2018). Unlike a formal Bayesian 3-D inversion, our framework targets process attribution of observed
123 variability and quantifies robustness through sensitivity tests rather than posterior flux estimates. This approach
124 emphasizes process-level interpretation of concentration variability while explicitly evaluating key uncertainty

sources. While CO is widely used to infer fossil-fuel CO₂, it carries inherent uncertainties due to varying emission ratios (R_{CO}), background variability, and atmospheric chemical processing (Turnbull et al., 2011; Chen et al., 2020; Griffin et al., 2007; Vimont et al., 2019). Footprint estimates are likewise sensitive to meteorological forcing and boundary-layer representation, and such transport sensitivities may be amplified under complex coastal mesoscale flows. We therefore quantify uncertainties associated with measurement and background selection, evaluate robustness to transport-model setup via STILT configuration tests, assess biospheric consistency using seasonal vegetation activity (e.g., NDVI), and utilize emission inventories for contextual comparison—using inter-inventory spread as a plausibility envelope for benchmarking rather than a validation target.

Accordingly, we aim to: (i) resolve spatiotemporal patterns of CO₂ across three sites spanning near-coastal (NS), urban-core (PY), and suburban (CH) environments; (ii) quantify how SLB modulates diurnal-seasonal CO₂ variability and interpret the contrasts in terms of changes in wind ventilation, boundary-layer depth, and seasonally varying atmospheric stability/stratification; and (iii) partition observed CO₂ enhancements at the urban site (PY) into fossil-fuel and biogenic components using CO and footprint information, with explicit uncertainty characterization via measurement/background terms and STILT configuration sensitivity tests.

2 Data and methodology

2.1 Observational sites

The high-precision greenhouse gas monitoring network in Guangzhou is illustrated in Fig. 1. Three stations—Nansha (NS), Panyu (PY), and Conghua (CH) are symmetrically distributed along the city’s predominant south–north wind axis, representing coastal, urban, and suburban atmospheric conditions, respectively. Site selection criteria are detailed in the Supplement. All stations employ tower-based sampling at similar heights: NS and PY at 48 m, and CH at 40 m. Monitoring spanned from January 1, 2023, to September 30, 2024. From PY, the straight-line distances to NS and CH are 54 km and 89 km, respectively.

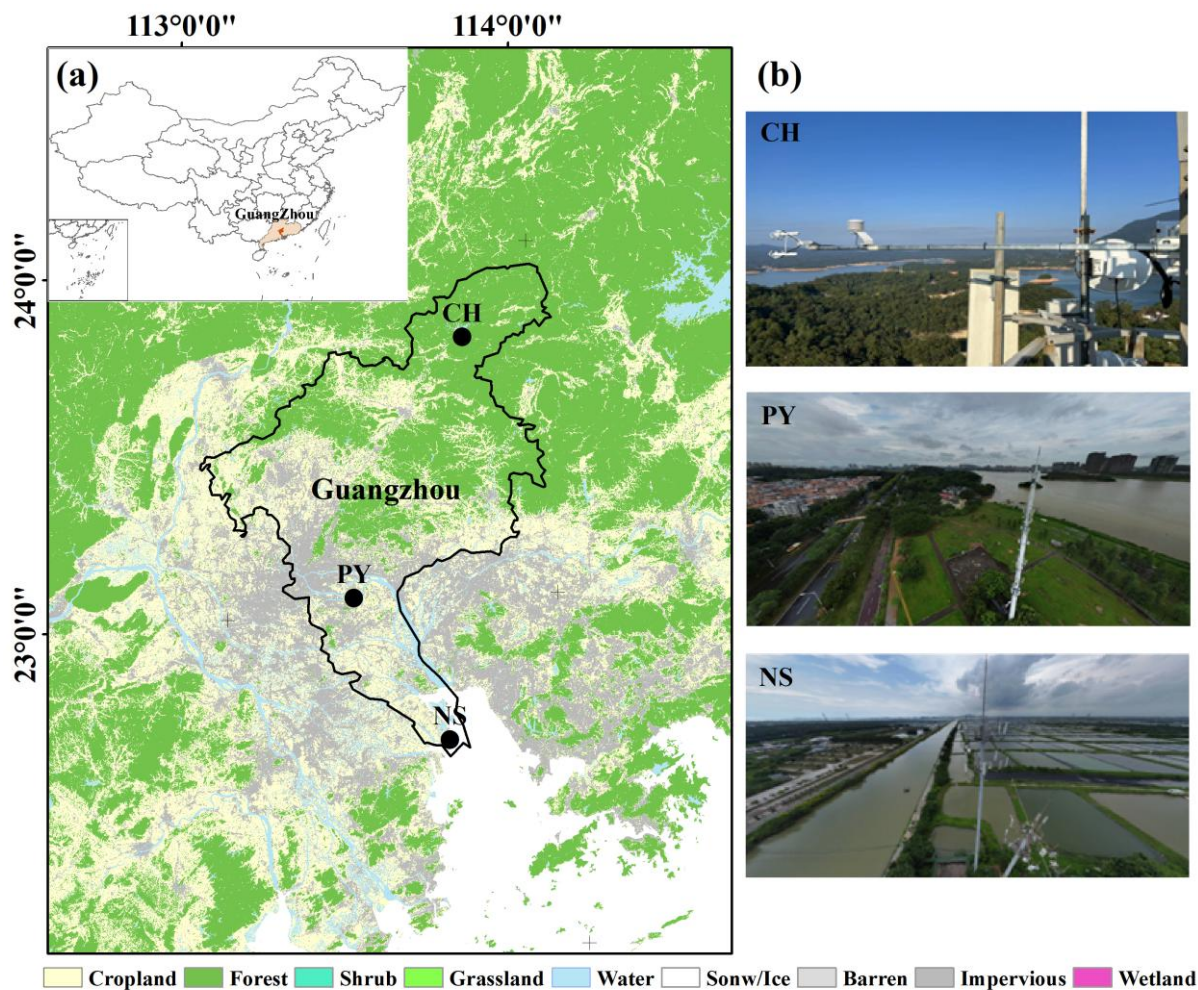


Figure 1. (a) Geographic locations of the NS, PY, and CH stations, with regional land use classification based on the 30 m resolution 2023 CLCD data (Yang and Huang, 2025). (b) Photograph of each station.

The NS station (113.63° E, 22.61° N) is located < 5 km from the coastline. This coastal site is surrounded by aquaculture ponds and sparse wetlands. Infrastructure nearby includes the under-construction southern extension of Guangzhou Metro Line 18 (NW direction) and the S78 highway (2 km west). The PY station (113.38° E, 23.03° N) is situated in the densely populated urban core; the tower is adjacent to Guangzhou University Town (north) and the Pearl River (south). A city road (100 m north) and the S73 expressway (700 m west) contribute to local emissions. The CH station (113.78° E, 23.74° N) is positioned in the northern suburbs. The site is bordered by subtropical evergreen broadleaf forests (north), a tourist resort (south), and a tea processing plant. The G45 highway lies 3 km northwest. According to the Emissions Database for Global Atmospheric Research (EDGAR) Community GHG database (EDGAR_2024_GHG; 2023; $0.1^{\circ} \times 0.1^{\circ}$) (Crippa et al., 2024), grid-level CO_2 emission densities for NS, PY, and CH are 3456, 15244, and 203 $\text{ton km}^{-2} \text{yr}^{-1}$, respectively (Fig. S1 in the Supplement).

2.2 Monitoring system

All three stations are equipped with similar monitoring systems, consisting of sampling modules, calibration

modules, gas analyzers, and data acquisition systems. Notably, the NS and PY stations utilize Picarro G2401 greenhouse gas analyzers to measure CO₂/CH₄/CO/H₂O, with a CO₂ measurement precision of < 20 ppb (5 min, 1 σ). The CH station employs an ABB GLA331-GGA greenhouse gas analyzer to measure CO₂/CH₄/H₂O, with a CO₂ measurement precision of < 25 ppb (5 min, 1 σ). N₂O and CO are measured using a GLA351-N2OCM analyzer. Detailed monitoring system and principles of the instruments are provided in the Supplement. Prior to field deployment, comparative tests were conducted in the laboratory to ensure the analytical performance consistency of the instruments. Additionally, meteorological sensors (measuring wind speed, direction, humidity, temperature, and pressure) are installed at the same height as the sampling inlets at NS and PY stations, while the CH station lacks such sensors. Detailed descriptions of wind field characteristics at NS and PY stations are included in the Supplement and illustrated in Fig. S2.

2.3 Calibration methods

The calibration module comprises two components: working standard curve establishment and target gas verification. High- and low-concentration standard gases are used to establish calibration curves, while a mid-concentration standard gas is used for target verification. The target and calibration gases are stored in inert-coated aluminum cylinders, uniformly supplied by the China National Environmental Monitoring Center. All stations follow the same calibration protocol: (1) weekly calibration curve establishment: high- and low-concentration gases are injected for 30 minutes each, with the final 5 minutes of instrument response used for calibration; (2) target gas verification every 12 h: mid-concentration gas is injected for 30 minutes, with the final 5 minutes of response used for verification; (3) re-calibration is triggered if the residual value (H) from target verification exceeds ± 0.2 ppm.

182

Calibration curves are derived from the instrument's response to calibration gas, yielding a linear calibration equation:

$$Y = A \times X - B, \quad (1)$$

where A, B are calibration coefficients. Calibrated CO₂ (CO_{2,k}) is calculated by:

$$CO_{2,k} = A \times CO_{2,m} - B, \quad (2)$$

where CO_{2,m} is the measured response. Daily 12 h target gas verification is conducted to assess analyzer accuracy and stability by calculating the residual H:

$$H = (A \times CO_{2,c} - B) - CO_{2,n}. \quad (3)$$

where CO_{2,n} is the standard CO₂ concentration of the target gas, and CO_{2,c} is the analyzer response/reading to the target gas.

194 To ensure high-precision and stable monitoring results, periods with $|H| \leq 0.1$ ppm are prioritized. Measurement
 195 uncertainties for the analyzers at NS, PY, and CH stations, calculated as the standard deviation (SD) of H (Yang et
 196 al., 2021), are 0.04, 0.02, and 0.04 ppm, respectively. In addition to daily calibration, maintenance personnel
 197 conduct weekly inspections of instruments and station facilities, including checks on power supply stability, data
 198 logger functionality, and industrial control computer status. Consumables (e.g., filters) are replaced as needed, and
 199 emergency repairs or instrument overhauls are performed when necessary. Any instrument downtime caused by
 200 internal or external factors is documented in maintenance logs, and affected data is flagged. Throughout the
 201 monitoring period, all three stations maintained data validity rates exceeding 90 %.

202 2.4 Sea–land breeze identification

203 The straight-line distances from the NS, PY, and CH stations to the coastline are 4, 58, and 130 km, respectively.
 204 The NS station, closest to the coast, was selected to study Guangzhou’s sea–land breeze (SLB) circulation. Prior to
 205 SLB identification, local and background winds must be differentiated, as tower-measured winds (Fig. S2 in the
 206 Supplement) represent superimposed local and background wind fields, where strong background winds can
 207 obscure SLB signals (Qiu and Fan, 2013). The following equations distinguish background winds from local winds
 208 (Sun et al., 2022; Shen et al., 2019):

$$209 \quad U_b = \overline{\sum_{i=0}^{23} U_i}, \quad (4)$$

$$210 \quad V_b = \overline{\sum_{i=0}^{23} V_i}, \quad (5)$$

$$211 \quad U_l = U_o - U_b, \quad (6)$$

$$212 \quad V_l = V_o - V_b. \quad (7)$$

213 where U_o and V_o are the observed wind fields from the tower, U_b and V_b denote background winds, and U_l and V_l
 214 represent local winds.

215

216 A sea–land breeze day (SLBD) is defined as any 24 h period exhibiting a distinct transition from sea breezes during
 217 the day to land breezes at night (Xiao et al., 2023). SLB identification criteria vary regionally due to differences in
 218 topography and coastline geometry (Huang et al., 2025). Guided by the SLB criteria in Supplementary Table S1
 219 and historical patterns for the Pearl River Estuary (Qiu and Fan, 2013; Zhang et al., 2024; Mai et al., 2024b), we
 220 define for the north-shore site (NS) directional sectors of $112\text{--}202^\circ$ for the sea breeze and $302\text{--}45^\circ$ for the land
 221 breeze. The corresponding detection windows are 12:00–20:00 LT (sea breeze) and 01:00–09:00 LT (land breeze),

222 consistent with regional SLB climatologies. Directional persistence is evaluated using the local-wind direction. The
 223 wind-speed screen is applied to the observed wind-speed magnitude at 48 m. A day is labeled as an SLB day only
 224 if all the following conditions are satisfied: (1) Directional persistence: within each window, the local wind
 225 direction (from the decomposed local component) stays inside the corresponding sector for ≥ 4 h, or for ≥ 4 h within
 226 any running 5 h window; and (2) Weak-forcing screen: for the same candidate SLB day, the observed wind-speed
 227 magnitude at 48 m must remain $< 10 \text{ m s}^{-1}$ at every hour of that day (i.e., no hourly value exceeds 10 m s^{-1}). This
 228 conservative cap follows coastal-China SLB climatologies and is intended to exclude strongly forced days (Qiu
 229 and Fan, 2013; Sun et al., 2022; Huang et al., 2025; Zhang et al., 2024). Otherwise, the day is designated a non-
 230 SLB day (NSLBD).

231
 232 Together, these two requirements reduce the likelihood of misclassification under strongly forced conditions. Days
 233 dominated by synoptic forcing or tropical-cyclone peripheries often show a prolonged anomalous wind regime
 234 rather than a clear diurnal reversal (Atkins and Wakimoto, 1997; Allouche et al., 2023). To explicitly assess residual
 235 tropical-cyclone (TC) contamination, we cross-referenced the SLB calendar with 2023 Pearl River Delta
 236 (PRD)/Guangzhou TC impact windows compiled from the official Guangdong–Hong Kong–Macao Greater Bay
 237 Area (GBA) Climate Monitoring Bulletin (Table S2). For each TC, Impact Start/End are defined as the first/last
 238 local dates on which the bulletin reports PRD/Guangzhou impacts or advisories attributable to that system
 239 (including peripheral rainbands/gusts). Because the bulletin is date-based, we conservatively treat the entire day
 240 within each window as potentially TC-influenced. Only one SLBD (2 Sep 2023) falls within these windows;
 241 excluding it leaves results unchanged.

242 **2.5 Estimation of CO_2tot , CO_2ff , and CO_2bio**

243 The observed CO_2 concentration enhancements at tall-tower receptor sites represent the integrated influence of
 244 upwind surface fluxes transported by atmospheric advection and mixing (Lin et al., 2003). Consequently, upwind
 245 carbon emissions can be inferred from site-specific enhancement measurements coupled with their corresponding
 246 atmospheric footprints. Here we apply an observation-driven framework that combines concentration-enhancement
 247 observations with transport-model footprints to estimate total CO_2 (CO_2tot) and CO (CO_{tot}) surface fluxes, and to
 248 further derive fossil-fuel (CO_2ff) and biogenic (CO_2bio) components using the site-specific $\Delta\text{CO}/\Delta\text{CO}_2$ relationship
 249 (R_{CO}), without relying on any a priori emissions inventory. The required inputs are limited to enhancement
 250 observations at the receptor sites and atmospheric footprints, consistent with previous regional flux quantifications
 251 for CO_2 , CH_4 , and CO (Mitchell et al., 2018; Lin et al., 2021; Wu et al., 2022a). Emission inventories are used only

for site-context characterization (Sect. 2.1) and, subsequently, for an independent bottom-up plausibility envelope for winter-afternoon flux estimates (Sect. 3.5), rather than as priors or validation targets. CO_{2tot} and CO_{tot} are calculated as:

$$CO_{2tot} = \frac{CO_{2,sobs} - CO_{2,bg}}{\sum_i Footprint_{i,s}}, \quad (8)$$

$$CO_{tot} = \frac{CO_{sobs} - CO_{bg}}{\sum_i Footprint_{i,s}}, \quad (9)$$

The numerators are hourly CO_2 and CO concentration enhancements (ΔCO_2 and ΔCO) at station s , where $CO_{2,sobs}$ and CO_{sobs} represent observed CO_2 and CO concentrations, while $CO_{2,bg}$ and CO_{bg} denote urban background concentrations (detailed in the Supplement). The denominators are hourly total atmospheric footprints ($\sum_i Footprint_{i,s}$), where i denotes backward particle release time from the receptor. Due to challenges in modeling mixed-layer depths during nighttime, morning, and evening, flux analysis focuses on afternoon hours (12:00–16:00) (Boon et al., 2016; Mitchell et al., 2018; Lin et al., 2021). Daily-scale CO_{2tot} and CO_{tot} are derived by dividing the mean afternoon ΔCO_2 and ΔCO by the corresponding mean $\sum_i Footprint_{i,s}$. The footprint quantifies the sensitivity of concentration enhancements at the observation site to upwind surface fluxes, as detailed in Sect. 2.5.1. ΔCO_2 is in units of [ppm], while footprint is in [ppm / ($\mu\text{mol m}^{-2} \text{s}^{-1}$)], so CO_{2tot} , the quotient between the two quantities, is in flux units of [$\mu\text{mol m}^{-2} \text{s}^{-1}$].

Anthropogenic CO_2 emissions (CO_{2ff}) are derived from CO_{tot} and the CO/ CO_2 emission ratio (R_{CO}), where R_{CO} is determined from real-time tower-measured data, as described in detail in Sect. 3.4:

$$CO_{2ff} = \frac{CO_{tot}}{R_{CO}}, \quad (10)$$

Biogenic fluxes (CO_{2bio}) are calculated as residuals:

$$CO_{2bio} = CO_{2tot} - CO_{2ff}. \quad (11)$$

Positive CO_{2bio} values indicate biogenic carbon emissions, while negative values denote carbon uptake, reflecting the dual role of urban biospheres as CO_2 sources and sinks (Kim et al., 2025).

2.5.1 Atmospheric transport model

To trace air mass sources entering the urban domain and reaching observation sites, and to assess CO_2 emissions corresponding to observed concentration enhancements, the Stochastic Time-Inverted Lagrangian Transport model (STILT-Rv2) was employed for atmospheric transport simulations, driven by meteorological fields from the Weather Research and Forecasting Model (WRFv4.1.1). In this study, STILT serves two purposes: (1) providing

280 airmass trajectories for identifying marine background concentrations in Guangzhou, and (2) generating
281 atmospheric footprints for quantifying total CO₂ and CO emissions.

282

283 The STILT model simulates atmospheric transport by releasing a set of air particles backward in time from the
284 receptor location at the observation height. These particles are tracked spatially and temporally as they disperse
285 upwind. The resulting trajectories delineate source regions influencing the receptor site and quantify the sensitivity
286 of observed concentrations to upwind surface fluxes, termed "source–receptor relationships" or "atmospheric
287 footprints" (Lin et al., 2003; Fasoli et al., 2018). Footprints represent the contribution of upwind sources/sinks to
288 downwind concentration changes, with higher sensitivities near receptors or under stable wind conditions, where
289 boundary layer airmasses interact more directly with surface fluxes (Wu et al., 2022a). For this study, 500 particles
290 were released at 48 m (PY station) heights, and traced backwards in time for 72 h. Footprints were computed at
291 $0.08^\circ \times 0.08^\circ$ spatial resolution. Periods with total footprint sensitivities ($\sum_i \text{Footprint}_i$) below the 10th percentile
292 were excluded, indicating low sensitivity to regional surface fluxes (Lin et al., 2021).

293

294 To evaluate whether STILT setup choices could bias the inferred fluxes, we performed targeted wintertime
295 sensitivity experiments at PY using paired daily comparisons ($n = 18$). Starting from the baseline (500 particles,
296 0.08° grid, 72 h backward), we independently varied (1) particle number (1000, 2000), (2) grid resolution (0.05° ,
297 0.10°), and (3) backward duration (96 h, 120 h). For each variant we recomputed footprints, reran the flux-
298 estimation framework, and compared paired daily afternoon means (12:00–16:00 LT) of CO₂tot, CO₂ff and CO₂bio
299 to the baseline using percent differences, Pearson r , and paired t -tests. Percent differences quantify effect size,
300 Pearson r assesses day-to-day consistency, and paired t -tests evaluate detectability of systematic mean shifts. This
301 paired-day design isolates transport-model parameter effects from day-to-day meteorology and observation
302 sampling.

303 2.5.2 Uncertainty sources

304 Uncertainties associated with the emission estimates inferred from Eqs. (12)–(14) primarily arise from four sources:
305 (1) observational uncertainty, (2) background concentration uncertainty, (3) atmospheric transport (footprint)
306 uncertainty, and (4) R_{CO}-related uncertainty. We do not explicitly propagate transport/footprint uncertainty within
307 the analytical error propagation of Eqs. (12)–(14); instead, we evaluate sensitivity to transport-model setup using a
308 winter paired-day STILT sensitivity analysis (Sect. 2.5.1; Sect. 3.5; Fig. 11). Residual transport biases (e.g., winds
309 and boundary-layer mixing) remain unquantified and may bias the inferred fluxes, thereby contributing to

inventory–observation differences when benchmarking against independent bottom-up inventories, alongside emission-inventory uncertainty and the representativeness mismatch between footprint-weighted enhancements and unweighted inventory means (discussed further in Sect. 3.5). Uncertainty associated with R_{CO} is not explicitly propagated as a formal regression-parameter uncertainty in this study; because R_{CO} is derived from CO and CO₂ enhancements, its uncertainty is expected to be driven mainly by uncertainties in the underlying enhancements (items 1–2) and scatter in the site-specific relationship. Among the quantified terms, uncertainties in CO₂tot are dominated by observational and background concentration errors. Uncertainties in CO₂ff are additionally affected by R_{CO} -related variability, while uncertainties in CO₂bio and the CO₂bio/CO₂ff offset ratio are propagated from CO₂tot and CO₂ff. Accordingly, the combined uncertainty term is calculated as:

$$E_u^2 = OBS_{u,c}^2 + BG_u^2, \quad (12)$$

where $OBS_{u,c}$ represents uncertainty in urban atmospheric observations, and BG_u represents uncertainty in urban background concentrations. We cannot accurately quantify all error sources involved in instrumental measurements; some minor error sources (e.g., uncertainty related to water vapor) may be negligible, while the primary uncertainty originates from discrepancies between measured concentrations and calibration standards (Verhulst et al., 2017). Here, $OBS_{u,c}$ is calculated as the standard deviation (SD) of residuals H (Yang et al., 2021). For urban background uncertainties:

$$BG_{u,co2}^2 = CT_{co2,r}^2 + CT_{co2,s}^2, \quad (13)$$

$$BG_{u,co}^2 = OBS_{co,r}^2 + OBS_{co,s}^2. \quad (14)$$

CO₂ background uncertainty ($BG_{u,co2}$) combines the absolute monthly smoothed residuals ($CT_{co2,r}$) and variability (SD) of monthly CO₂ concentrations ($CT_{co2,s}$) from CarbonTracker (CT). Similarly, CO background uncertainty ($BG_{u,co}$) is derived from monthly smoothed residuals ($OBS_{co,r}$) and variability ($OBS_{co,s}$) of in situ observations.

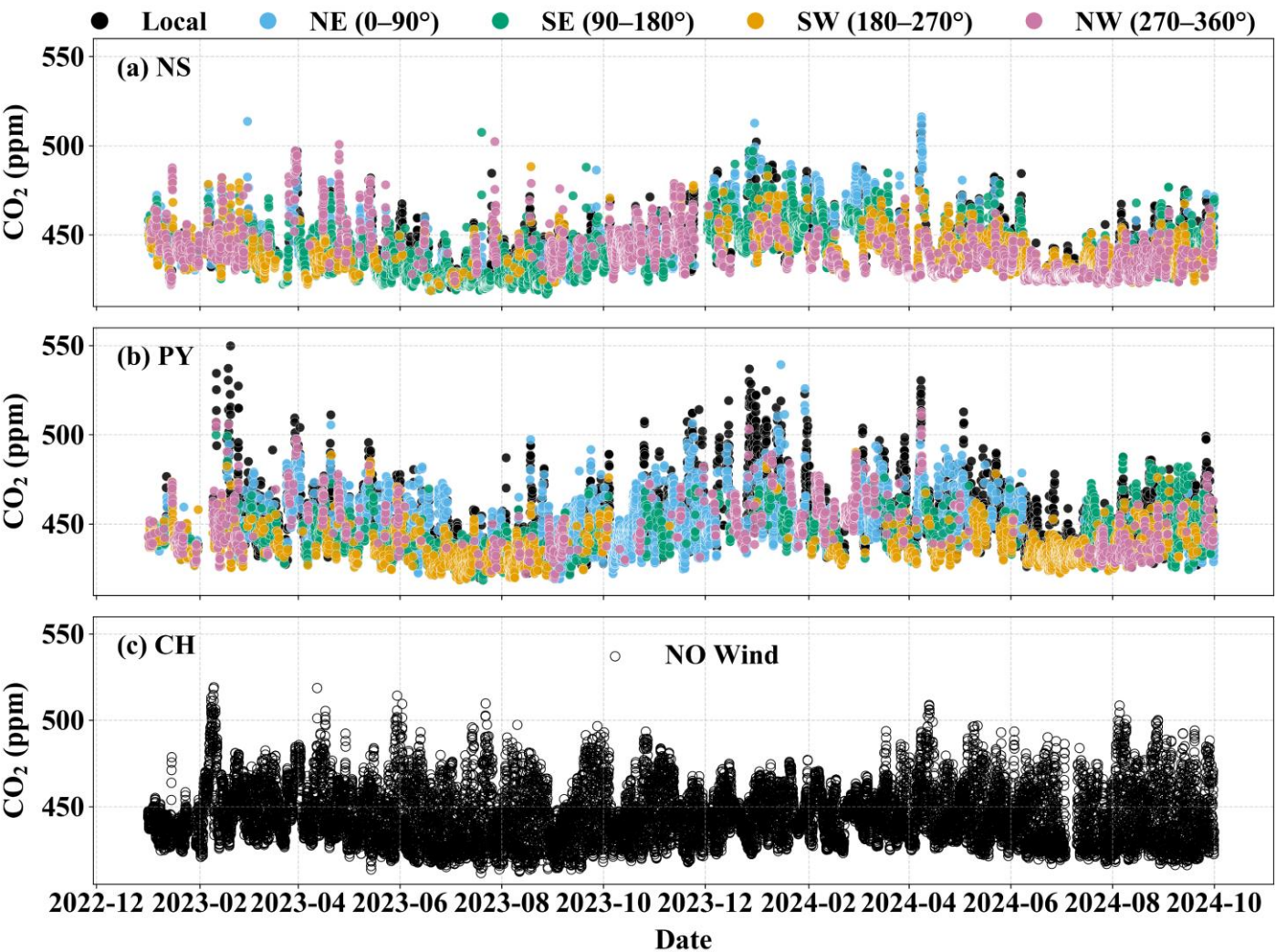
To guard against bias from transport settings, the paired-day sensitivity analysis is conducted relative to the baseline; we treat effect sizes and 95 % confidence intervals (CIs) as primary quantities and report p-values only to indicate detectability given $n = 18$, with quantitative outcomes presented in Sect. 3.5 and Fig. 11.

3 Results and discussion

3.1 Spatiotemporal patterns of atmospheric CO₂

Figure 2 presents the hourly mean time series of atmospheric CO₂ concentrations at the NS, PY, and CH stations

339 in Guangzhou from January 1, 2023, to September 30, 2024. To assess wind field impacts on CO₂ variability (Fig.
 340 S2), concentrations were classified into five categories: local (wind speed < 1.5 m s⁻¹) and four directional sectors
 341 (wind speed ≥ 1.5 m s⁻¹) defined as NE (0–90°), SE (90–180°), SW (180–270°), and NW (270–360°). All stations
 342 exhibited significant temporal variability, with standard deviations (SD) of 13.90 (NS), 15.92 (PY), and 16.05 ppm
 343 (CH), consistent with urban and suburban observations in Hangzhou, Beijing, Xi'an, and Seoul (Park et al., 2021;
 344 Yang et al., 2021; Chen et al., 2024; Liu et al., 2025). PY showed the highest CO₂ levels and variability, driven
 345 predominantly by local-type emissions under low wind speeds. In contrast, the NS coastal station exhibited
 346 elevated concentrations during northerly winds (NW/NE). Seasonal wind effects were pronounced:
 347 summer southerly winds (SW/SE) generally reduced CO₂ at PY and NS (most notably at NS near the coast), while
 348 winter northerly winds (NW/NE) often increased CO₂ at NS.



349
 350 **Figure 2.** Time series of atmospheric CO₂ concentrations at the (a) NS, (b) PY, and (c) CH stations. For NS and PY,
 351 points are color-coded by wind category: local conditions (wind speed < 1.5 m s⁻¹) and four directional sectors for winds ≥
 352 1.5 m s⁻¹ (NE, 0–90°; SE, 90–180°; SW, 180–270°; NW, 270–360°). For CH, wind-direction classification is not shown and
 353 the time series is plotted without sector coloring.

Urban–rural CO₂ gradients vary globally due to differences in economic activity, population density, land use, and energy infrastructure, reflecting heterogeneous urban carbon emissions (Gao et al., 2022). To further illustrate this spatial contrast and its seasonality, we summarized the urban–suburban/coastal gradients across the full record and report annual and seasonal means in Fig. 3. In Guangzhou, mean CO₂ concentration differences between PY and NS/CH were 6.67 and 3.43 ppm, respectively, forming a distinct "urban dome" (urban > suburban > coastal). The NS–CH difference (3.44 ppm) highlights comparable gradients between suburban and coastal zones. This gradient mirrors Los Angeles’s coastal megacity profile but with a smaller magnitude (Verhulst et al., 2017). Guangzhou’s urban–suburban difference (3.43 ppm) aligns with Hangzhou’s 2021 observations (4.96 ppm) (Chen et al., 2024) but is lower than Nanjing (8.1 ppm, 2014) and Beijing (12.4 ppm, 2018–2019) (Gao et al., 2018; Yang et al., 2021). It remains far smaller than Shanghai (55.1 ppm, 2014) and Baltimore (66.0 ppm, 2002–2006) (Pan et al., 2016; George et al., 2007). Over time, urban emissions may stabilize as suburban populations and fossil fuel demand grow, potentially narrowing urban–suburban CO₂ differences (Mitchell et al., 2018). For instance, Hangzhou’s reduced gradient reflects urbanization-driven energy consumption, where suburban monitoring captures urban emission influences (Chen et al., 2024).

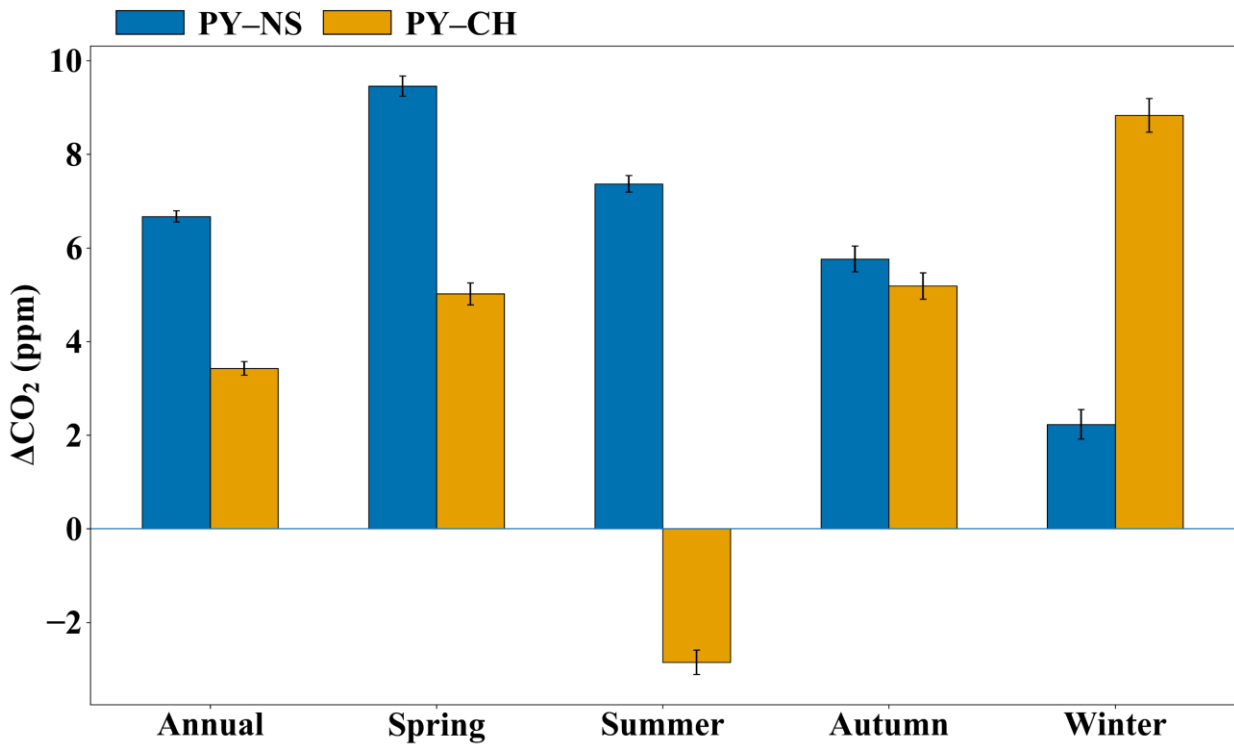


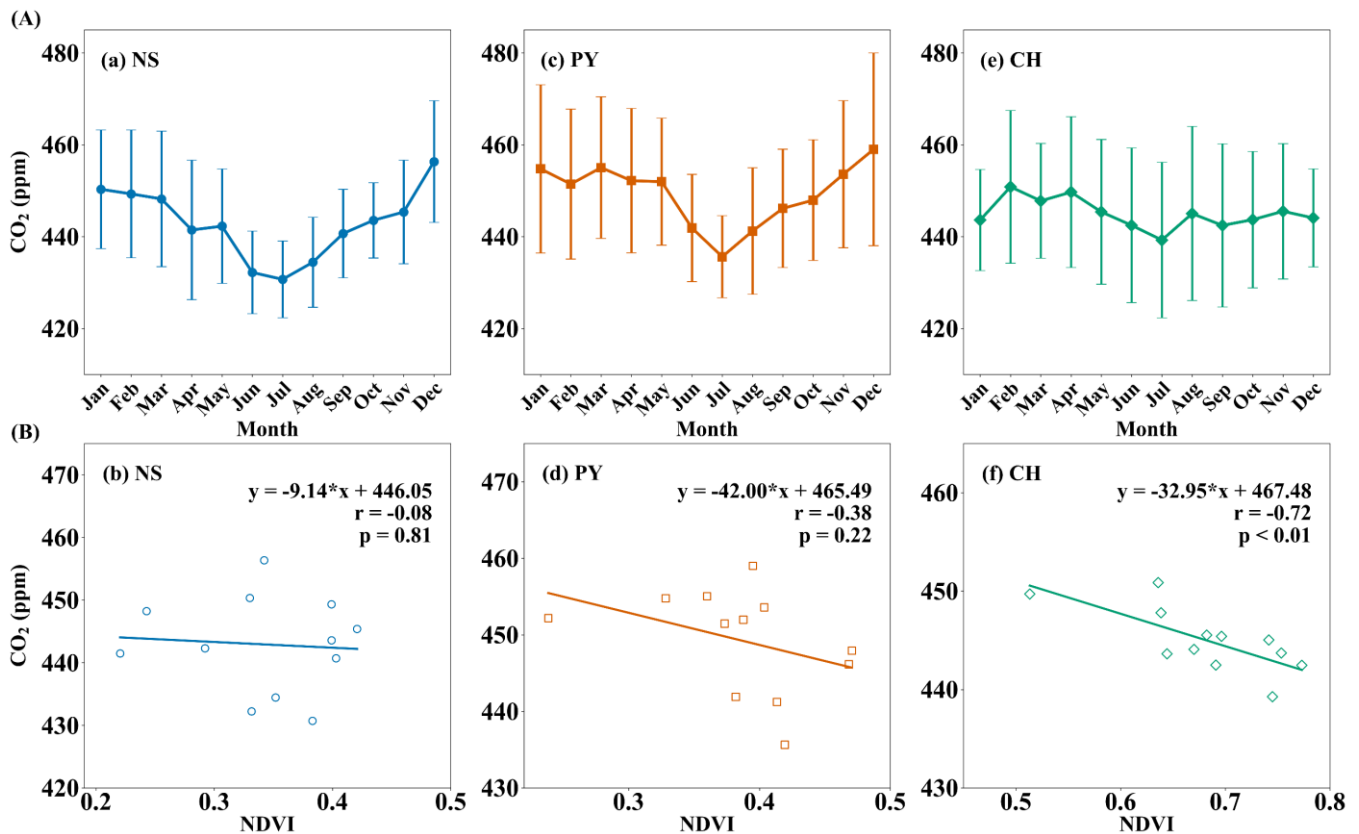
Figure 3. Urban–suburban/coastal CO₂ gradients in Guangzhou. Annual and seasonal mean concentration differences (ΔCO_2 , ppm) between the urban site (PY) and the coastal site (NS) (PY–NS) and between PY and the suburban site (CH) (PY–CH). Seasons are defined as Spring (Mar–May), Summer (Jun–Aug), Autumn (Sep–Nov), and Winter (Dec–Feb). Error bars denote ± 1 standard error (SE).

Figure 3 further shows that these spatial gradients are strongly season-dependent. PY–NS remains positive year-

round but is smallest in winter (2.23 ppm), when prevailing northerlies elevate CO₂ at the coastal NS site and reduce the urban–coastal contrast, consistent with Fig. 2. PY–CH is more strongly seasonally modulated: it peaks in winter (8.83 ppm) but reverses sign in summer (–2.86 ppm), when southerly (marine-influenced) flow more frequently ventilates PY (SW/SE sectors in Fig. 2) and the CH summertime enhancement may reflect a northward-displaced urban influence plus biogenic/boundary-layer modulation (Sect. 3.1.1; Sect. 3.3). Spring and autumn show intermediate positive PY–CH (~5 ppm), while PY–NS peaks in spring (9.46 ppm), consistent with enhanced coastal ventilation under southerly influence. Together, these gradients indicate a seasonally displaced CO₂ “dome” in this coastal setting, where the highest CO₂ within our network can occur outside the urban core, complementing the site-level seasonal cycles discussed in Sect. 3.1.1. This coastal, seasonally displaced CO₂ dome contrasts with patterns more commonly reported in inland megacity networks, where enhancements are typically strongest at the most urbanized sites relative to suburban/background stations (Xueref-Remy et al., 2018; Yang et al., 2021; Chen et al., 2024).

3.1.1 Seasonal variability of atmospheric CO₂

Figure 4 illustrates the monthly mean variations in atmospheric CO₂ concentrations at the NS, PY, and CH stations in Guangzhou, alongside their correlations with the Normalized Difference Vegetation Index (NDVI). NDVI data at 1 km × 1 km spatial resolution were obtained from NASA’s EOSDIS Land Processes Distributed Active Archive Center (Didan, 2015), with values within a 3 km radius buffer around each station center used for comparative analysis. All three stations exhibited consistent seasonal CO₂ patterns, with higher concentrations in winter/spring and lower values in summer/autumn, mirroring observations in Hangzhou (Chen et al., 2024). These variations arise from the combined effects of (1) seasonal biogenic flux cycles, (2) anthropogenic emission variability, and (3) boundary layer height dynamics (Xueref-Remy et al., 2018). Enhanced vegetation photosynthesis during warmer months (summer/autumn, Table S3 in the Supplement) strengthens biogenic carbon sinks, while higher boundary layer depths (Fig. S6 in the Supplement) and southerly marine air masses (Fig. 2) promote atmospheric mixing and CO₂ dispersion.



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Figure 4. (A) Variations in monthly mean CO₂ concentrations and (B) their correlations with Normalized Difference Vegetation Index (NDVI) for the (a–b) NS, (c–d) PY, and (e–f) CH stations. Error bars indicate ± 1 standard deviation (SD). The amplitudes of the seasonal variation of CO₂ at NS, PY, and CH are 25.63, 23.38, and 11.59 ppm, respectively. NS and PY peaked in December and troughed in July, whereas CH peaked in February and troughed in July. NS’s large amplitude reflects its extreme December highs and July lows. In December, prevailing northerly winds (Figs. 2a and S2a) transported urban emissions to downwind NS, narrowing its CO₂ difference with PY to 2.68 ppm. Conversely, July saw NS’s CO₂ concentrations fall to the lowest among all stations—4.93 ppm and 8.56 ppm below PY and CH, respectively—establishing a south-to-north increasing gradient (coastal < urban < suburban). This gradient aligns with marine-influenced southerly air masses, which dilute coastal CO₂ while transporting urban emissions northward, potentially accumulating CO₂ in northern suburbs. This south–north pattern is consistent with the seasonal mean gradients in Fig. 3, which summarize how the urban–coastal contrast weakens in winter and how the urban–suburban gradient can be seasonally displaced in summer. Although CH exhibits strong biogenic coupling (NDVI–CO₂ correlation: –0.72; Fig. 4f), NS shows an opposite seasonal contrast relative to CH (–9.80 ppm in summer; +5.80 ppm in winter), pointing to transport- and boundary-layer-driven variability at the coastal site. At NS, the NDVI–CO₂ correlation is weak (–0.08) and NDVI varies within a narrow range (0.22–0.42), indicating limited local biogenic control, whereas seasonal shifts in marine–

continental transport (summer dilution vs. winter urban outflow) and boundary-layer depth jointly modulate dilution and accumulation (Figs. 2 and S6). In February, CH recorded its annual CO₂ maximum, driven by vegetation respiration during early growth stages and elevated emissions from fireworks around the Lunar New Year, as CH's location falls outside Guangzhou's fireworks restriction zones (https://www.gz.gov.cn/gfxwj/qjgfxwj/chq/qf/content/post_7198980.html, last access: 18 June 2025). This interpretation is supported by CO, a tracer for combustion-derived CO₂ (Newman et al., 2013; Che et al., 2022a): CH's CO concentrations also peaked in February due to firework emissions, whereas other sites peaked in December (Fig. S7 in the Supplement).

3.1.2 Diurnal variations of atmospheric CO₂

The diurnal patterns of atmospheric CO₂ concentrations at NS, PY, and CH stations in Guangzhou consistently exhibited lower daytime and higher nighttime values (Fig. 5). This is attributed to the shallow nocturnal boundary layer, which traps anthropogenic and biogenic fluxes near the surface, elevating CO₂ levels. After sunrise, surface heating deepens the boundary layer, diluting surface emissions and entraining free tropospheric air with lower CO₂ concentrations. Concurrently, daytime photosynthetic uptake further reduces near-surface CO₂ (Mitchell et al., 2018). We further evaluate urban–suburban–coastal differences in these processes. At PY, the CO₂ peak occurred at 08:00–09:00, aligning with morning traffic peaks, reflecting dominant anthropogenic influences. CH's peak appeared 1–2 h earlier than PY due to its longitudinal and elevational position, where earlier sunrise accelerates the breakup of the nocturnal stable boundary layer. Both PY and CH reached minima at 16:00–17:00, likely linked to afternoon photosynthetic activity. NS exhibited irregular peak/valley timing.

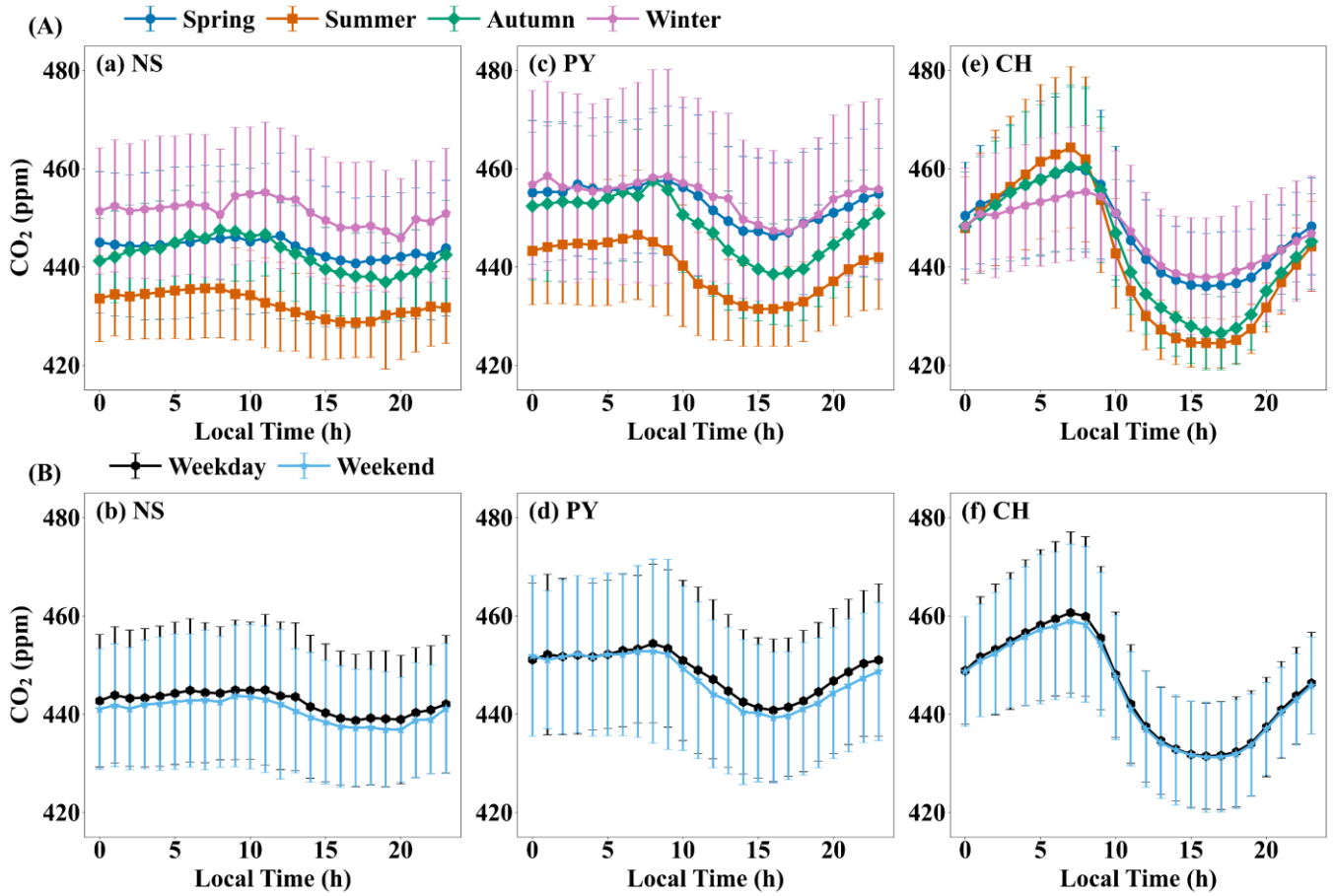


Figure 5. Diurnal CO₂ variations at the (a–b) NS, (c–d) PY, and (e–f) CH stations across (A) seasons and (B) weekdays/weekends. Seasons are defined as Spring (Mar–May), Summer (Jun–Aug), Autumn (Sep–Nov), and Winter (Dec–Feb). Error bars indicate ± 1 SD. The corresponding CO diurnal cycles are shown in Fig. S9.

Diurnal amplitudes at CH and PY were larger in summer/autumn than winter/spring, driven by vegetation activity and boundary layer dynamics (Fig. S8 in the Supplement). Summer/autumn conditions in Guangzhou—abundant light, warmth, and rainfall—optimize vegetation growth, enhancing daytime photosynthesis and nighttime respiration (Dusenge et al., 2019). Optimal canopy temperatures for subtropical evergreen forests ($\sim 30^\circ\text{C}$) (Liu et al., 2015) align with CH/PY's summer/autumn daytime temperatures (Table S3), explaining their amplified amplitudes. However, the diurnal amplitude of CO₂ at CH in summer and autumn is 2.63 times and 1.77 times that at PY, respectively. The diurnal amplitude of atmospheric CO₂ concentration at CH in summer is 39.90 ppm, which is close to the diurnal amplitude of CO₂ concentration in the suburbs of Hangzhou in summer (35.29 ppm) (Chen et al., 2024). Despite similar temperatures, CH's larger NDVI range and stronger NDVI-CO₂ correlation (-0.72 vs. PY; Figs. 4d and f) highlight greater biogenic dominance, with pronounced daytime uptake and nighttime respiration. NS showed the smallest diurnal amplitudes across seasons (e.g., 5.60 ppm in summer), attributable to sparse vegetation (low NDVI: 0.22–0.42) and frequent summer southerly marine air masses, which dilute coastal CO₂.

Figure 5B contrasts weekday–weekend diurnal CO₂ patterns. All stations generally showed higher weekday concentrations, diverging from Hangzhou and Beijing (Yang et al., 2021; Chen et al., 2024) but aligning with Paris and Boston (Briber et al., 2013; Xueref-Remy et al., 2018). At CH, the daytime weekday–weekend contrast is small. This could reflect a weak anthropogenic weekday–weekend signal relative to other sources of variability, but it does not uniquely diagnose source dominance because transport and boundary-layer mixing may dilute or mask weekday–weekend differences. At PY, the clearer daytime weekend decrease is consistent with reduced on-road activity in the urban core. At NS, weekday CO₂ remains higher across much of the day, which may reflect weekday-intensified construction and port-related logistics in the surrounding area (for example, the Metro Line 18 extension and operations near the Nansha Container Terminal Phase III, ~5 km east), superimposed on transport and mixing effects.

To corroborate the combustion contribution to the CO₂ diurnal cycle, we examined synchronous CO measurements (Fig. S9 in the Supplement). At all sites, CO shows a consistent seasonal ordering (winter highest, summer lowest), and its morning maximum at PY (typically 08:00–10:00) coincides with the CO₂ morning peak (Fig. 5), supporting traffic/combustion control of the early-day enhancement. In contrast to CO₂, CO exhibits only a weak mid-afternoon minimum (including at CH), consistent with CO₂ being additionally depressed by photosynthetic uptake while CO is unaffected; both species remain modulated by transport and boundary-layer mixing. Weekday–weekend differences are small at CH but clearer at PY/NS (Fig. S9), indicating a stronger anthropogenic weekly-cycle imprint in the urban-core and coastal/port settings. Overall, the CO–CO₂ contrast reinforces our interpretation that the morning CO₂ maxima are primarily combustion-driven, whereas the pronounced mid-afternoon CO₂ minima at vegetated sites reflect biogenic uptake rather than reduced emissions.

3.2 Sea–land breeze impacts

Based on meteorological observations from the NS coastal tall tower, 84 sea–land breeze days (SLBD) were identified in Guangzhou between January 2023 and September 2024, accounting for 13.14 % of the monitoring period, with peaks in spring and autumn. These transitional seasons between summer and winter are characterized by weaker synoptic systems and lighter background winds, favoring SLBD occurrence (Mai et al., 2024b). Our results align with SLBD seasonal distributions for the Pearl River Estuary cities of Zhuhai and Guangzhou in 2022 (Zhang et al., 2024; Mai et al., 2024b). Figure 6 compares CO₂ concentrations during SLBD and non-SLB days (NSLBD) across stations. Overall, average CO₂ concentrations during SLBD were lower than during NSLBD by 5.87 ppm at NS, 3.08 ppm at PY, and 0.75 ppm at CH. This indicates that SLB circulation enhances ventilation and

483 CO₂ dispersion, with the SLBD-related reduction decreasing from the coastal site to the urban core and being
484 smallest at the suburban site (NS > PY > CH). This pattern is similar to SLB-driven dispersion reported for PM_{2.5},
485 PM₁₀, and ozone in Tianjin (Hao et al., 2017). Seasonal differences were pronounced: SLB promoted CO₂
486 dispersion at NS and PY in spring, winter, and autumn (spring > winter > autumn), but increased CO₂ accumulation
487 in summer. Tianjin similarly observed summer PM_{2.5}/PM₁₀ accumulation under SLB (Hao et al., 2017). At CH,
488 SLBD reduced CO₂ in spring, summer, and autumn but increased it in winter, likely due to limited inland SLB
489 penetration and competing winter processes.

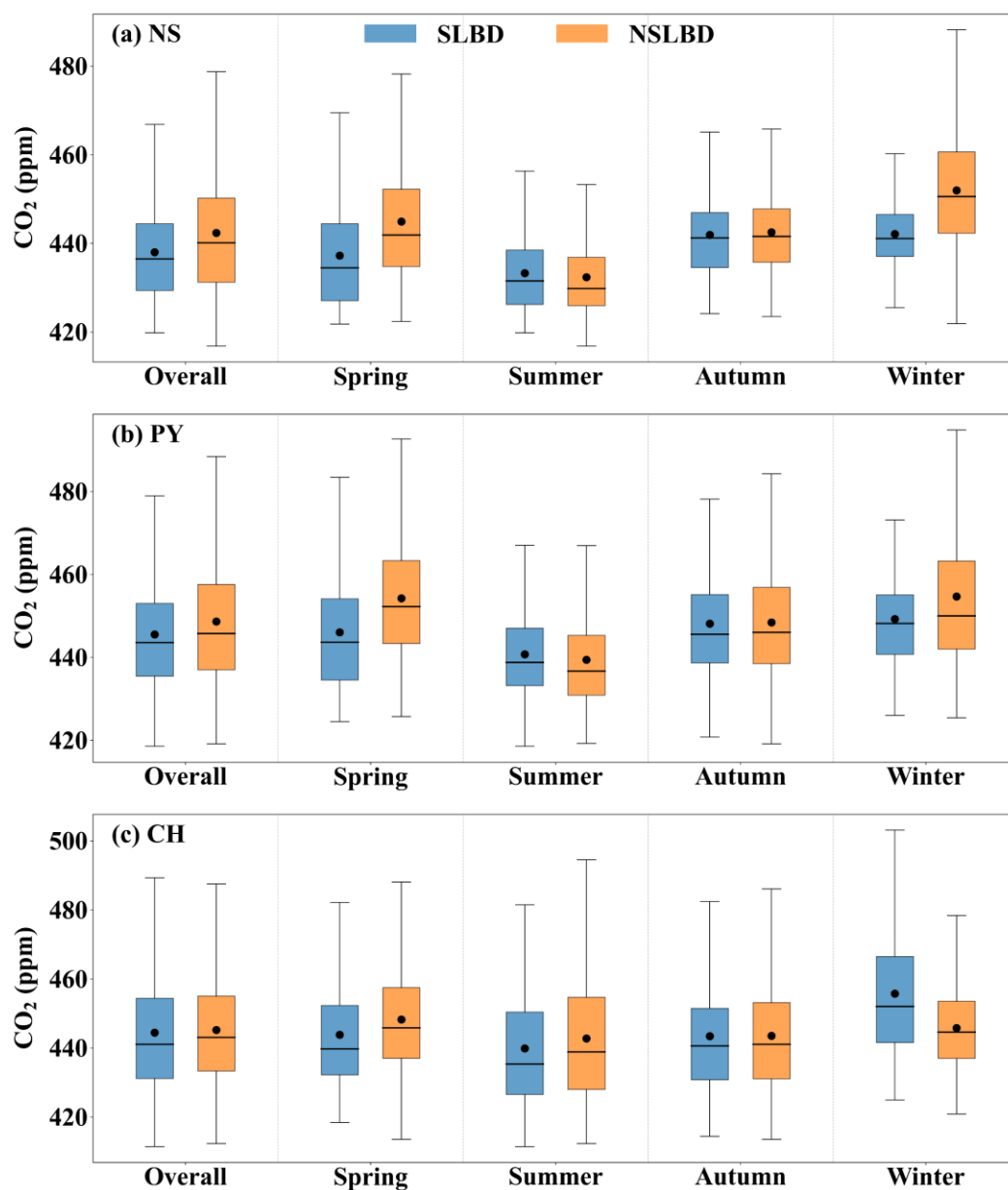
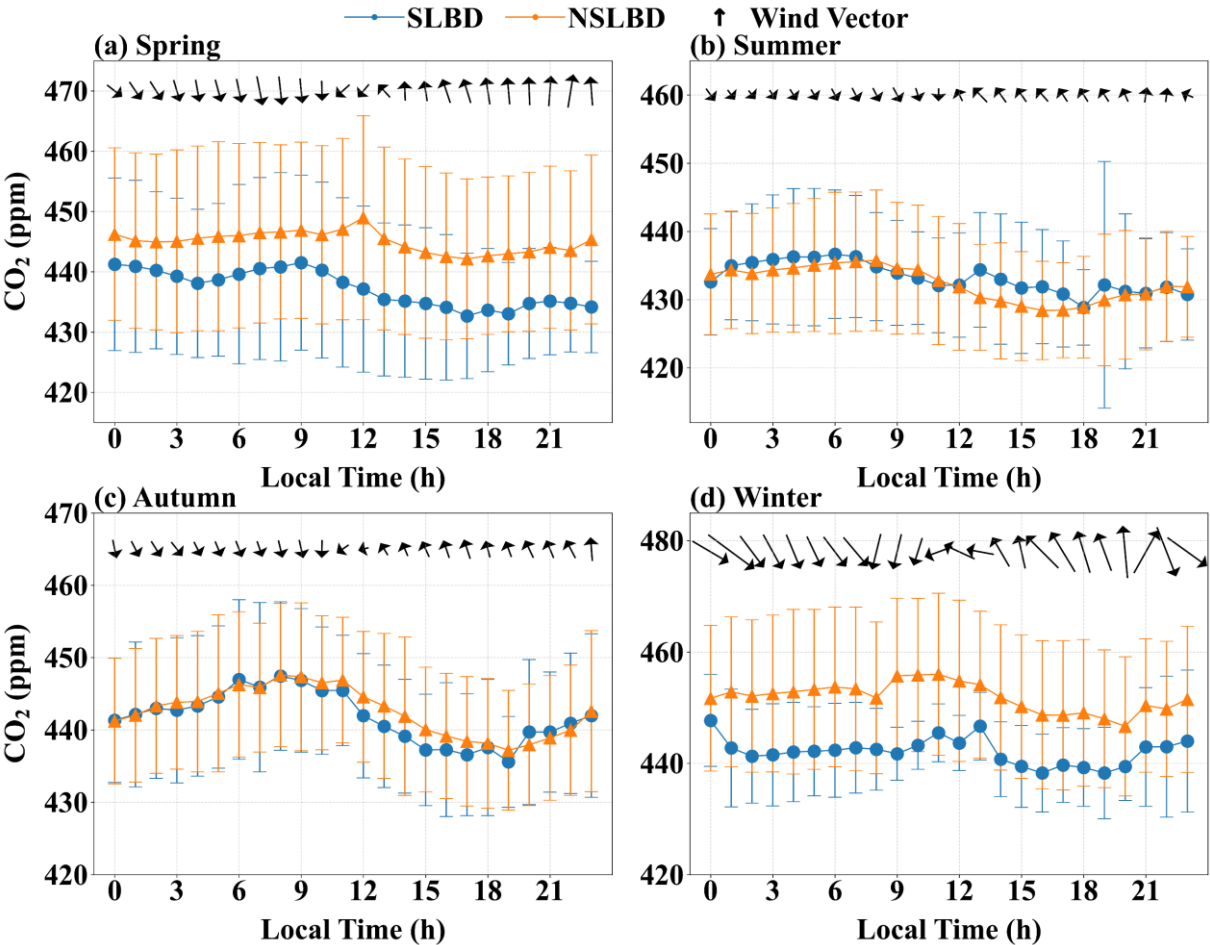


Figure 6. Boxplots of atmospheric CO₂ concentrations (black dots denote means; black line denotes the median) during sea-land breeze days (SLBD) and non-SLB days (NSLBD) by station and season, with outliers excluded.

493 To resolve seasonal and diurnal SLB impacts, we analyzed CO₂ diurnal variations during SLBD and NSLBD (Fig.
494 7). Focusing on NS (due to similar PY–NS trends and space constraints), spring and winter SLBD reduced CO₂

495 concentrations by 7.76 ppm and 9.77 ppm (hourly mean differences), respectively, driven by stronger winds (Fig.
 496 7) and deeper boundary layers (Fig. S10 in the Supplement). Autumn SLB only reduced CO₂ during sea breeze
 497 hours (mean difference: 1.69 ppm). Autumn's weaker winds and boundary layers resulted in reduced dispersion
 498 compared to spring/winter. In summer, SLB increased CO₂ by 2.08 ppm (sea breeze hours) due to stable
 499 atmospheric stratification. Summer temperatures were 6.00 °C and 12.19 °C higher than spring and winter (Table
 500 S3), respectively. Under calm, rain-free conditions, the collision of moist marine air with dry-hot coastal land
 501 formed a thermal internal boundary layer (TIBL), inducing low-level temperature inversions near the SLB
 502 convergence zone (Liu et al., 2001; Reddy et al., 2021). These inversions suppressed horizontal/vertical mixing,
 503 trapping CO₂ (Stauffer et al., 2015; Hao et al., 2024). NS's summer SLBD winds averaged 1.05 m s⁻¹ (sea breeze)
 504 and 0.96 m s⁻¹ (land breeze)—38.60 %, 63.16 %, and 15.32 % lower than spring, winter, and autumn winds,
 505 respectively—while boundary layer heights (590.54 m) were 9.51 % shallower than NSLBD (Fig. S10). Weak
 506 winds and shallow boundary layers stabilized atmospheric stratification, limiting CO₂ dispersion and elevating
 507 ground-level CO₂ by up to 4.03 ppm.



508
 509 **Figure 7.** Diurnal variations in CO₂ concentrations, wind direction, and wind speed at the coastal station (NS) during
 510 sea-land breeze days (SLBD) and non-SLB days (NSLBD) by season. Error bars indicate ± 1 SD. The corresponding CO
 511 diurnal cycles are shown in Fig. S11.

CO provides a combustion-specific tracer to further diagnose SLB modulation of anthropogenic signals (Fig. S11 in the Supplement). In spring and winter, CO is generally lower on SLBD than on NSLBD, consistent with enhanced dilution and export by the breeze circulation, broadly mirroring the CO₂ behavior (Fig. 7). In autumn, the CO response is weaker and transitional, with only modest daytime reductions. In summer, however, SLB days exhibit a pronounced afternoon CO build-up, with a much larger relative enhancement than CO₂ (relative to the seasonal 24 h mean on NSLBD: ~10 % for CO versus ~0.7 % for CO₂), implying trapping/recirculation of fresh combustion plumes under weak-wind, shallow-mixing conditions. Overall, the joint CO–CO₂ behavior confirms that SLB exerts a seasonally varying control on near-surface carbon signals—ventilating in cooler seasons but favoring accumulation under the stagnant summer regime.

3.3 CO₂ enhancements and uncertainties

Figure S12 (in the Supplement) presents the time series of observed CO₂ and CO concentrations at Guangzhou's stations relative to marine backgrounds from January 1 to December 27, 2023. Compared to urban observations with significant hourly variability, marine background concentrations in Guangzhou remained stable, with summer and winter CO₂ standard deviations of 0.94 ppm and 0.67 ppm, respectively, indicating minimal local source/sink influences. Using Eqs. (13) and (14), marine background uncertainties were calculated (Table S4 in the Supplement). Summer and winter CO₂ marine background uncertainties were 0.96 ppm and 0.70 ppm, respectively, constraining urban marine background uncertainties below 1 ppm—slightly lower than Los Angeles's 1.4 ppm (Verhulst et al., 2017). CO marine background uncertainties were 12.68 ppb (summer) and 18.36 ppb (winter).

Based on background concentrations, CO₂ enhancements were derived for all stations. Figure 8 shows enhancements across all hours, afternoon (12:00–16:00), and midnight (00:00–05:00) periods in 2023, summer, and winter. Annual median enhancements were 13.59 (NS), 17.70 (PY), and 16.29 ppm (CH), with pronounced spatiotemporal variability—closely aligning with the 10–20 ppm range observed annually in the Beijing-Tianjin-Hebei (BTH) urban cluster of China (Han et al., 2024). In summer, the all-hours enhancement followed a south-to-north gradient: 7.00 (NS), 13.23 (PY), and 16.91 ppm (CH). This inland maximum likely reflects the combined influence of coastal transport, biogenic exchange, and boundary-layer mixing, and is consistent with the seasonal gradient displacement discussed in Sect. 3.1 (Fig. 3). Afternoon enhancements peaked at PY (6.92 ppm), whereas midnight enhancements at CH reached 31.36 ppm. Winter afternoon enhancements reversed this pattern: 16.58 (NS), 12.37 (PY), and 7.45 ppm (CH), with NS and PY values 4.39 times and 1.79 times higher than summer. Midnight enhancements at CH remained highest in winter (18.87 ppm), despite a 38.93 % reduction from summer.

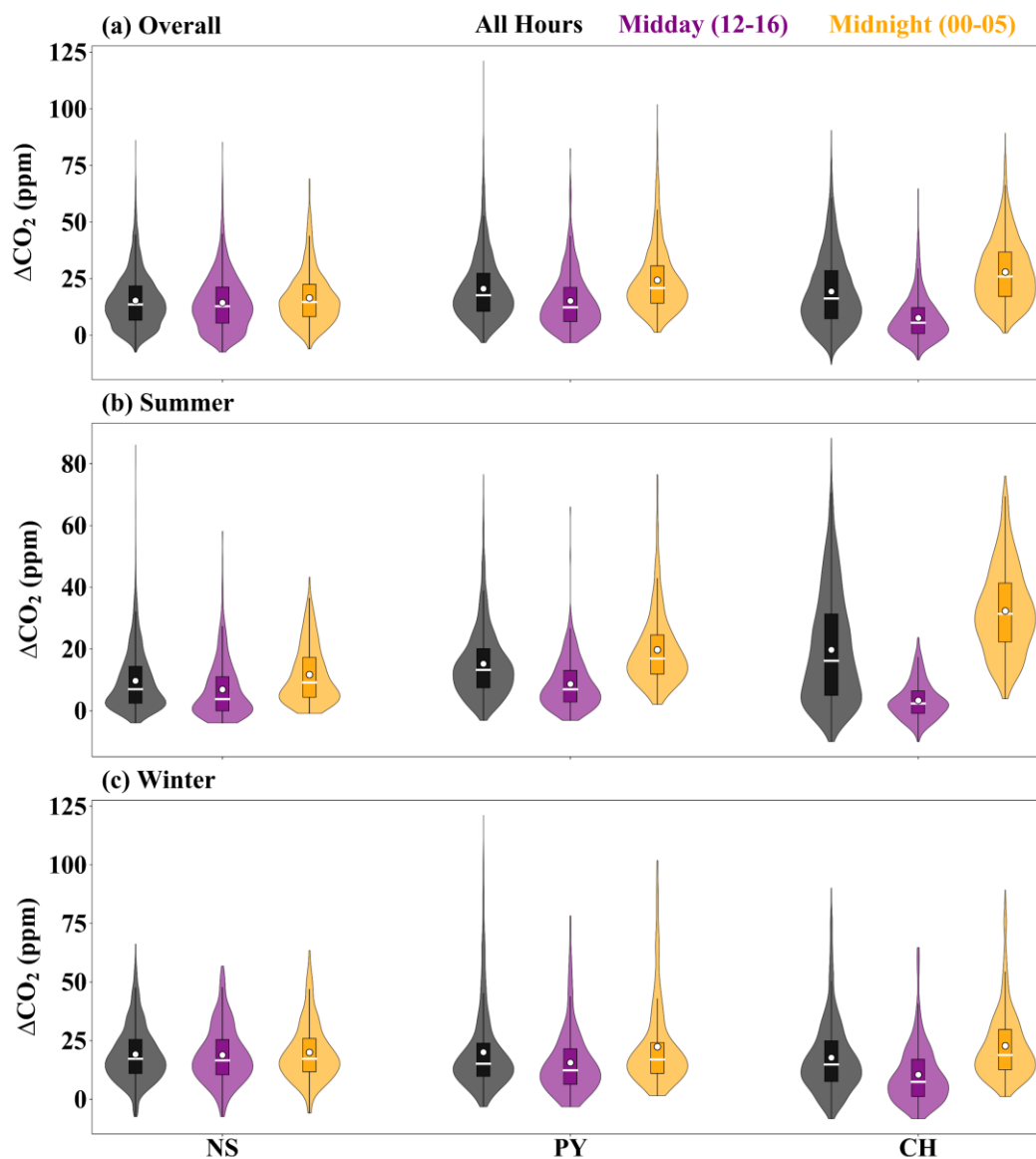


Figure 8. Distributions of hourly CO₂ enhancement (ΔCO_2) above the background concentrations at NS, PY, and CH during the (a) overall, (b) summer, and (c) winter periods, shown for all hours, midday (12:00–16:00 LT), and midnight (00:00–05:00 LT). White dots denote the mean values, and white horizontal lines denote the median values.

This spatiotemporal variability reflects divergent influences of anthropogenic emissions, biogenic fluxes, and atmospheric mixing. At CH, strong diurnal shifts in enhancements (e.g., 31.36 ppm summer midnight) highlight biogenic dominance, with long-tailed distributions (Fig. 8). Stable, shallow nighttime boundary layers trapped respiratory emissions near the surface, consistent with isotopic studies in Xi'an (32.80 ppm) and Switzerland (30.00 ppm) (Wang et al., 2021; Berhanu et al., 2017). At NS, transport dominated: summer southerly marine air masses reduced enhancements, while winter northerly winds transported urban emissions downstream, raising NS enhancements to PY levels (exceeding PY in afternoons). PY's enhancements were primarily anthropogenic, validated by CO co-variation. CO, a tracer for combustion-derived CO₂, showed significantly higher concentrations at PY (Fig. S12). PY's median midnight CO enhancements in summer were 2.04 times and 1.43 times higher than

NS and CH (Fig. S13 in the Supplement). Shallow nocturnal boundary layers localized anthropogenic CO near the surface, with minimal vertical/horizontal transport, confirming PY's anthropogenic dominance.

3.4 Continuous observations of $\Delta\text{CO}/\Delta\text{CO}_2$ ratios

Reduced Major Axis regression (Model II) was applied to analyze the relationship between CO (ΔCO) and CO_2 (ΔCO_2) concentration enhancements across stations, with the $\Delta\text{CO}/\Delta\text{CO}_2$ emission ratio (R_{CO}) derived from regression slopes (Fig. 9). In 2023, R_{CO} values for NS, PY, and CH were 8.48 ± 1.81 , 7.45 ± 1.38 , and 4.16 ± 3.59 ppb ppm⁻¹, respectively, with correlation coefficients of 0.78, 0.72, and 0.33, indicating significant spatiotemporal heterogeneity. Summer R_{CO} was generally lower than winter, with CH exhibiting the lowest seasonal value (2.66 ± 0.83 ppb ppm⁻¹). Winter maxima occurred at NS (11.03 ± 1.49 ppb ppm⁻¹), followed by PY (9.08 ± 1.13 ppb ppm⁻¹) and CH (8.56 ± 1.82 ppb ppm⁻¹).

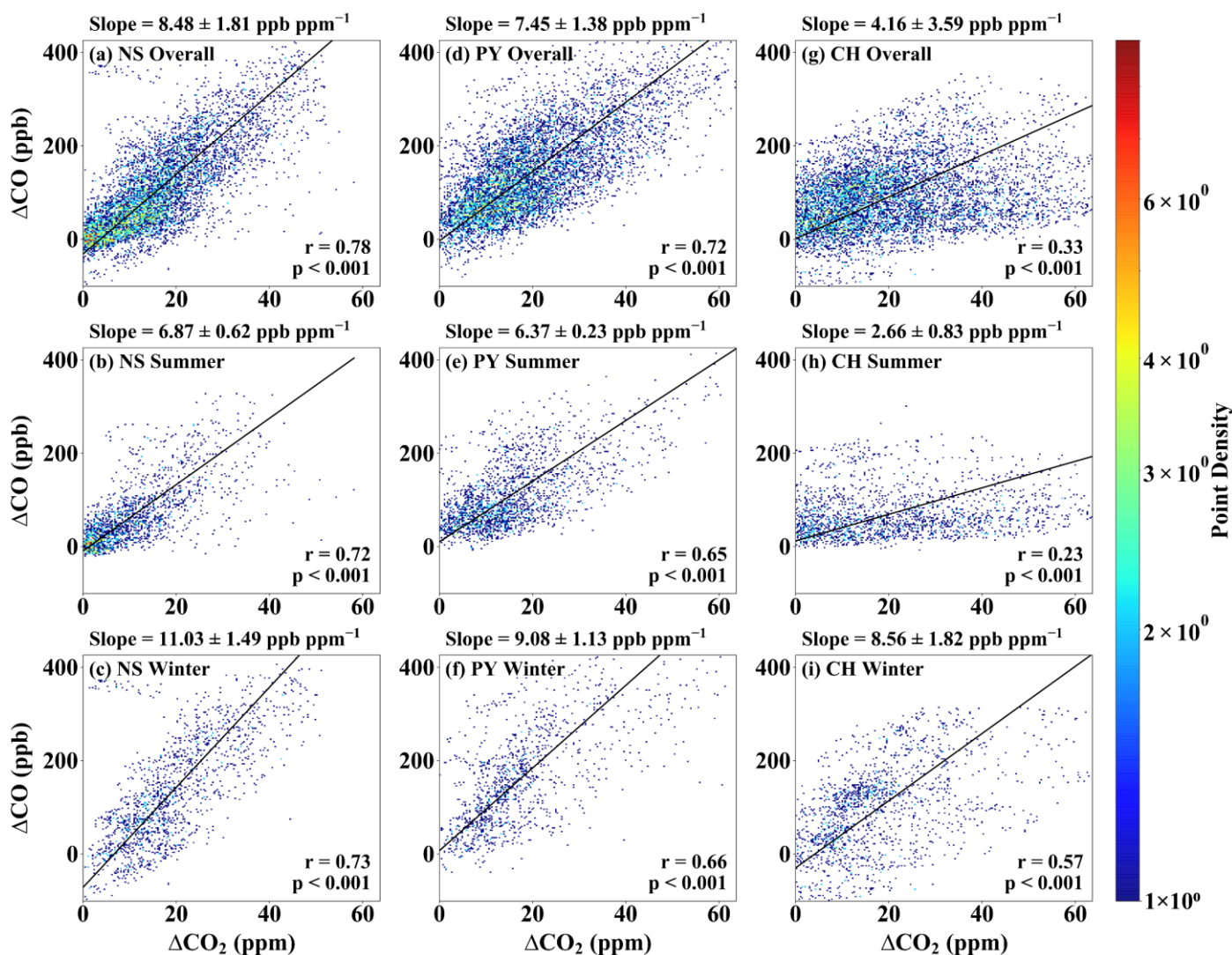


Figure 9. Seasonal relationships between ΔCO_2 and ΔCO enhancements at the (a–c) NS, (d–f) PY, and (g–i) CH stations, analyzed using geometric-mean regression. Panels are shown as 2D histogram density plots (hist2d; 200 × 200 bins), where color indicates the number of paired observations per bin. The fitted slope represents the $\Delta\text{CO}/\Delta\text{CO}_2$ emission ratio (R_{CO} ; ppb ppm⁻¹), reported as mean ± 1 SD (reflecting temporal variability).

Comparatively, Beijing's urban R_{CO} in 2019 was measured at 10.46 ± 0.11 ppb ppm⁻¹ using portable Fourier-transform spectroscopy (Che et al., 2022a), while Shanghai and Los Angeles showed 10.22 ± 0.40 and 9.64 ± 0.46 ppb ppm⁻¹, respectively, based on satellite and model data (Wu et al., 2022a). Guangzhou's lower R_{CO} is consistent with the post-2013 tightening of China's air-quality management across the energy and transport sectors, rather than a single intervention. National action plans (2013–2017 Air Pollution Prevention and Control Action Plan; 2018–2020 Three-Year “Blue-Sky” Action Plan) strengthened coal/industrial and vehicle-emission controls with explicit targets and timelines. In parallel, ultra-low-emission (ULE) retrofits of coal-fired power units were rolled out through 2020, and China 6 (VI) on-road emission standards were phased in, with large cities (including Guangzhou) leading implementation; Guangdong's provincial Blue-Sky measures further reinforced industrial/mobile-source controls and promoted fleet electrification. These measures coincide with independent inventory evidence of declining national CO emissions (~23 % during 2013–2017) (Zheng et al., 2018), plausibly reducing CO/CO₂ emission ratios from dominant urban sources. Consistently, Beijing's R_{CO} decreased from > 30 ppb ppm⁻¹ in 2006 (Han et al., 2009) to 10.22 ± 0.40 ppb ppm⁻¹ by 2020 (Wu et al., 2022a), with additional reductions during the 2008 Olympics and 2020 COVID-19 lockdowns (Wang et al., 2010; Cai et al., 2021). In Guangdong, restrictions on coal plants, retirement of inefficient industries, and promotion of electric vehicles coincided with large declines in SO₂ and NO₂ (85 % and 35 % in 2019 relative to 2006) (Hu et al., 2021), and Mai et al. (2021) likewise reported improved combustion efficiency in the PRD associated with advances in gasoline vehicles. We emphasize that this interpretation is consistency-based rather than a formal causal identification; fuel mix, fleet composition, and atmospheric oxidation may also contribute (Young et al., 2023; Vimont et al., 2019).

Seasonal R_{CO} variations stem from biogenic exchange and transport dynamics. Summer's weaker ΔCO – ΔCO_2 correlations at CH reflect dominant biogenic influences (daytime uptake and nighttime respiration), as reported in Beijing, Indianapolis, and Switzerland (Turnbull et al., 2015; Berhanu et al., 2017; Che et al., 2022a). Biogenic impacts decreased from suburban > urban > coastal, aligning with vegetation gradients. Winter's higher R_{CO} at CH and NS correlated with reduced biogenic activity and northerly transport of urban emissions under stable boundary layers. Berhanu et al. (2017) attributed winter R_{CO} increases to cold-air advection and boundary layer accumulation. NS's winter R_{CO} (4.16 ppb ppm⁻¹ higher than summer) linked to urban airmass origins, while PY's seasonal shifts reflected suburban source–sink variations. Although secondary CO from upwind Volatile Organic Compounds (VOCs) and CH₄ oxidation could perturb R_{CO} , their combined contribution was merely 1 % in coastal urban regions (Griffin et al., 2007).

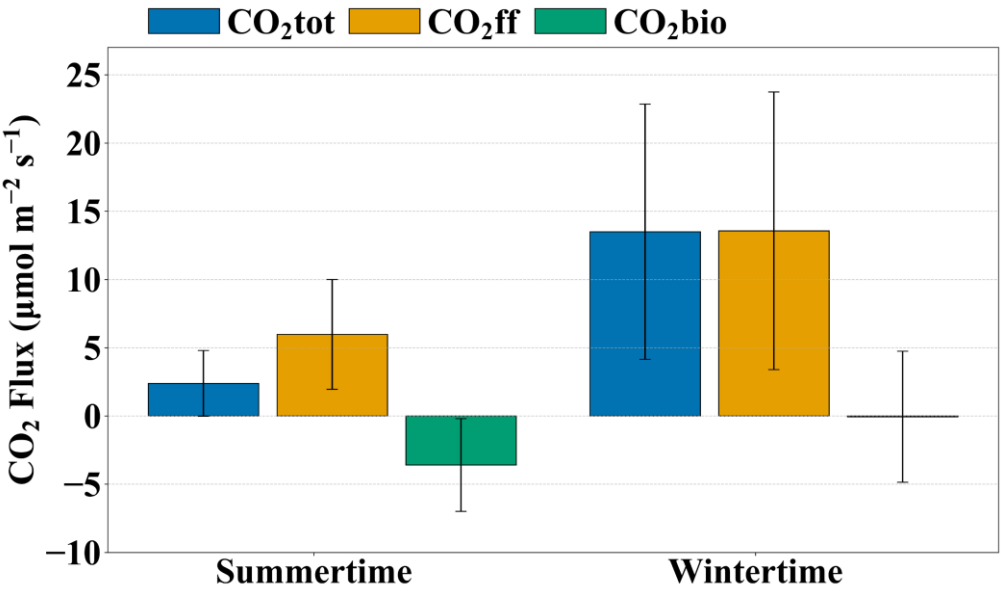
600 3.5 Partitioning anthropogenic and biogenic fluxes

601 Section 3.4 shows that the sites differ in how well the $\Delta\text{CO}-\Delta\text{CO}_2$ relationship reflects fossil-fuel combustion. At
602 CH, especially in summer, CO_2 variability is strongly influenced by biogenic exchange, which weakens the
603 combustion linkage implied by R_{CO} and can bias an R_{CO} -based fossil-fuel estimate. At NS, in contrast, variability
604 is dominated by changing transport and upwind airmass origin, particularly in winter when urban plumes frequently
605 reach the coastal site. We therefore select PY—the urban-core site with a comparatively robust combustion signal—
606 to quantify the surface CO_2 emissions ($\text{CO}_{2\text{tot}}$) constrained by the observed concentration enhancements and
607 footprint-informed transport. Using the PY-specific R_{CO} , we then partition $\text{CO}_{2\text{tot}}$ into fossil-fuel ($\text{CO}_{2\text{ff}}$) and
608 biogenic ($\text{CO}_{2\text{bio}}$) components.

609

610 Figure 10 summarizes mean afternoon (12:00–16:00 LT) $\text{CO}_{2\text{tot}}$, $\text{CO}_{2\text{ff}}$, and $\text{CO}_{2\text{bio}}$ at PY for July 2023 (summer)
611 and December 2023 (winter). Bars show monthly means of daily afternoon values. The plotted error bars indicate
612 ± 1 standard deviation (SD) across days and thus represent day-to-day variability in ventilation, mixing, and
613 transport rather than uncertainty of the monthly mean. Mean uncertainty is quantified by the standard error (SE),
614 which is distinct from the SD shown in Fig. 10. In July, $\text{CO}_{2\text{tot}} = 2.38 \pm 0.45$, $\text{CO}_{2\text{ff}} = 5.97 \pm 0.75$, and $\text{CO}_{2\text{bio}} =$
615 $-3.59 \pm 0.63 \mu\text{mol m}^{-2} \text{s}^{-1}$ (mean $\pm$ SE), whereas in December $\text{CO}_{2\text{tot}} = 13.50 \pm 2.20$, $\text{CO}_{2\text{ff}} = 13.56 \pm 2.40$, and
616 $\text{CO}_{2\text{bio}} = -0.06 \pm 1.13 \mu\text{mol m}^{-2} \text{s}^{-1}$ (Table S5). Because $\text{CO}_{2\text{bio}}$ is diagnosed as a residual ($\text{CO}_{2\text{bio}} = \text{CO}_{2\text{tot}} -$
617 $\text{CO}_{2\text{ff}}$), its uncertainty reflects propagated uncertainties from both $\text{CO}_{2\text{tot}}$ and $\text{CO}_{2\text{ff}}$ —including measurement and
618 background-selection uncertainty and R_{CO} -related variability—rather than being independent. Notably, December
619 $\text{CO}_{2\text{bio}}$ is close to zero. The bootstrap 95 % CI of the monthly mean is $[-2.28, 2.09] \mu\text{mol m}^{-2} \text{s}^{-1}$, which includes
620 zero, indicating that the net biogenic flux was not statistically distinguishable from zero during winter afternoons,
621 likely reflecting a near-neutral balance between photosynthesis and respiration. In contrast, the July mean $\text{CO}_{2\text{bio}}$
622 remains clearly negative relative to its uncertainty, supporting robust summertime net biogenic uptake despite
623 uncertainty propagation inherent to the residual calculation. When assessed using daily afternoon means, the July–
624 December contrasts are statistically significant for $\text{CO}_{2\text{tot}}$, $\text{CO}_{2\text{ff}}$, and $\text{CO}_{2\text{bio}}$ (Welch and Mann–Whitney tests;
625 bootstrap 95 % confidence intervals; Table S5). Robust distributional metrics corroborate this significant seasonal
626 increase despite partial day-to-day overlap: the $\text{CO}_{2\text{ff}}$ median increases from 4.33 (IQR: 3.58–7.81) in July to 10.70
627 (IQR: 6.74–18.28) $\mu\text{mol m}^{-2} \text{s}^{-1}$ in December. Across both months, $\text{CO}_{2\text{ff}}$ is larger in magnitude than $\text{CO}_{2\text{bio}}$,
628 implying that the PY afternoon enhancement is primarily fossil-fuel driven, consistent with fossil-dominated urban
629 enhancements reported for other Chinese cities (Wang et al., 2022). The stronger winter $\text{CO}_{2\text{ff}}$ relative to summer
630 is explained mainly by atmospheric dynamics—reduced dilution under weaker marine ventilation and a shallower

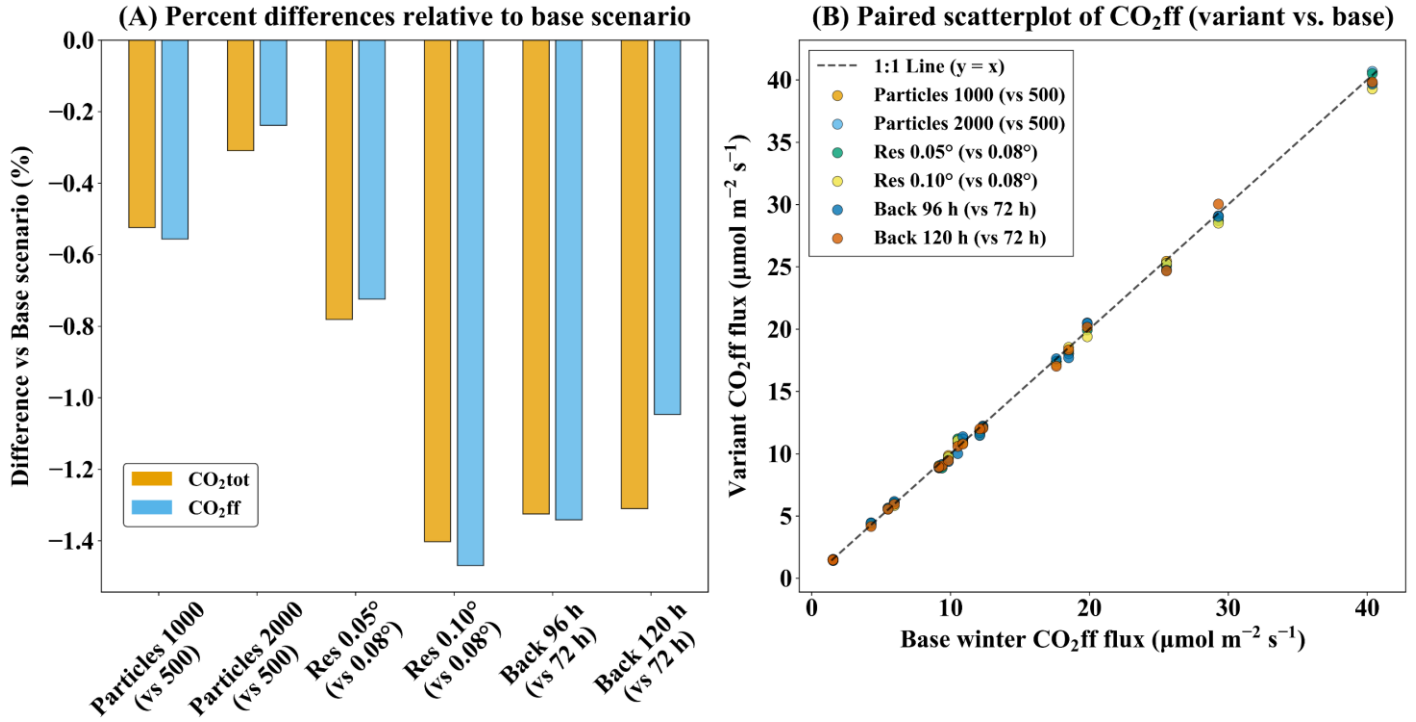
631 boundary layer—while seasonal emission changes (e.g., winter residential energy use) likely provide a secondary
 632 contribution. In contrast, CO₂bio is more negative in summer, consistent with higher summer NDVI (+11 %) and
 633 warmer conditions approaching optimal canopy temperatures (Table S3).



634 **Figure 10.** Average afternoon (12:00–16:00 LT) CO₂tot, CO₂ff, and CO₂bio at PY for July 2023 (summer; n = 29 valid
 635 days) and December 2023 (winter; n = 18 valid days). December has fewer valid days because objective QC excluded days
 636 with incomplete afternoon coverage (e.g., instrument downtime/maintenance), so the smaller winter sample reflects data
 637 availability rather than subjective selection. Bars show monthly means of daily afternoon values. Error bars indicate ± 1
 638 standard deviation (SD) across daily afternoon means within each month (day-to-day atmospheric variability), not the
 639 standard error (SE) of the monthly mean; SE and confidence intervals are reported in Table S5.

641 To test the robustness of the winter fossil-fuel dominance inferred at PY—when combustion signals are strongest—
 642 we reran the winter flux-estimation workflow using paired daily afternoon means (12:00–16:00 LT). We then varied
 643 (1) particle number (1000/2000), (2) grid spacing (0.05°/0.10°), and (3) backward duration (96/120 h) relative to
 644 the baseline (500 particles, 0.08°, 72 h). Mean changes were small for both components (Table S6 in the Supplement;
 645 Fig. 11A). Increasing particle number to 1000/2000 changed CO₂ff by −0.56 %/−0.24 % and CO₂tot by
 646 −0.52 %/−0.31 %. Refining the grid to 0.05° yielded comparably small decreases (CO₂ff: −0.72 %; CO₂tot:
 647 −0.78 %). Extending the backward duration to 96/120 h produced changes of −1.34 %/−1.05 % for CO₂ff and
 648 −1.33 %/−1.31 % for CO₂tot. Only the coarser 0.10° grid produced a statistically detectable, yet small, decrease
 649 (CO₂ff: −1.47 %, p = 0.0269; CO₂tot: −1.40 %, p = 0.0164). All other settings yielded changes ≤1.34 % with 95 %
 650 CIs spanning zero (p ≥ 0.083), indicating no evidence of material bias at our sample size. Day-to-day consistency
 651 remained essentially unchanged across settings, with extremely high correlations (CO₂ff: r = 0.9993–0.9997;
 652 CO₂tot: r = 0.9992–0.9997) and small RMSE values (CO₂ff: 0.31–0.45 μmol m⁻² s⁻¹; CO₂tot: 0.28–0.45 μmol m⁻²
 653 s⁻¹), consistent with the near-1:1 paired scatter (Fig. 11B). Because wintertime CO₂bio at PY was close to zero,

percent differences were not informative; in absolute terms, test–baseline differences in CO₂bio remained small (order 10⁻² μmol m⁻² s⁻¹) with 95 % CIs generally spanning zero, consistent with the tight across-run daily spread (median 0.045 μmol m⁻² s⁻¹; IQR 0.016–0.067 μmol m⁻² s⁻¹) (Table S7 in the Supplement). Across the baseline plus six variants, the day-by-day ensemble spread was tightly bounded (median 0.20–0.21 μmol m⁻² s⁻¹, median CV ≈ 1.8 %). Together, these results indicate that our baseline STILT configuration lies in a converged regime and that the inferred winter CO₂ff dominance is robust to reasonable transport-parameter choices.

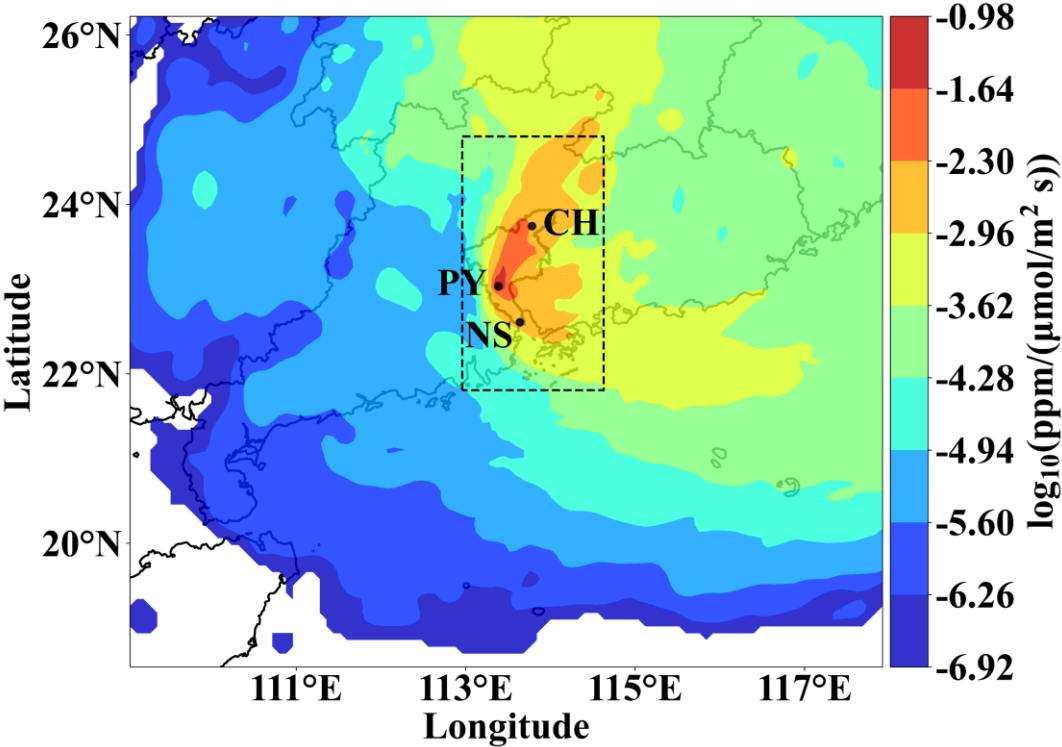


660

Figure 11. STILT parameter sensitivity at PY (winter). Panel A: mean percent difference (variant – baseline) of inferred fluxes relative to the winter baseline (500 particles, 0.08°, 72 h), computed from paired daily afternoon means (12:00–16:00 LT; n = 18); Δ% = (variant – base)/base × 100; negative values indicate lower than baseline. Panel B: paired scatter of CO₂ff (μmol m⁻² s⁻¹) from each variant versus the baseline for the same days; solid line is 1:1 (y = x).

For winter afternoons at PY, measurement and background-selection uncertainties—estimated by propagating their combined enhancement uncertainty through Eqs. (12)–(14)—contribute only ~0.36 μmol m⁻² s⁻¹, i.e., ~3 % of the mean winter-afternoon CO₂ff. Paired-day STILT sensitivity tests (Fig. 11) further indicate that the winter-afternoon CO₂ff attribution is robust to reasonable transport-parameter choices. The remaining uncertainty is thus more likely dominated by residual transport representation error and coastal mesoscale flow (e.g., wind and boundary-layer mixing biases), as documented for winter conditions in meteorological/transport modeling (Yadav et al., 2021; Lin et al., 2021). To provide independent context, we benchmark our winter-afternoon estimate against bottom-up inventories, which are used solely for contextual comparison and do not enter the CO₂ partitioning. We use 2023 inventories at 0.1° × 0.1° resolution, temporally disaggregate annual totals to hourly emissions using the same profiles (Crippa et al., 2020), and compute the winter-afternoon (12:00–16:00 LT) mean emission flux over the

675 winter footprint-defined sensitivity region (Fig. 12). This yields $19.81 \mu\text{mol m}^{-2} \text{s}^{-1}$ from the EDGAR_2024_GHG
 676 (2023) inventory (Crippa et al., 2024) and $85.46 \mu\text{mol m}^{-2} \text{s}^{-1}$ from the MEIC-global-CO₂ (2023) inventory (Xu et
 677 al., 2024). While this aligns qualitatively with higher MEIC estimates reported for Guangdong Province (Yang et
 678 al., 2025), the contrast is larger for our winter-afternoon sensitivity region. This likely reflects differences in spatial
 679 disaggregation and point-source representation (e.g., road-network allocation and power-plant locations) that can
 680 be accentuated at sub-provincial scales (Yang et al., 2025).



681
 682 **Figure 12.** Spatial distribution of the mean winter-afternoon (12:00–16:00 LT) STILT footprint at the PY station,
 683 representing the receptor sensitivity to upwind surface fluxes. Colors denote the log₁₀-transformed footprint sensitivity (ppm
 684 per (μmol m⁻² s⁻¹)). The dashed rectangle outlines the sensitivity region used for inventory flux aggregation and
 685 benchmarking in Sect. 3.5.

686 Notably, our observation-based CO₂ff reflects the footprint-weighted enhancement sampled at PY during winter
 687 afternoons after background removal, whereas inventories provide gridded emission fields from which an
 688 unweighted domain-mean afternoon flux can be computed over the sensitivity region. Because only grid cells with
 689 substantial STILT footprint influence contribute materially to the receptor enhancement—and these weights are
 690 highly heterogeneous and transport-dependent (Fig. 12)—the unweighted domain-mean inventory flux may be non-
 691 uniformly represented in the receptor signal under variable coastal transport (Gerbig et al., 2003; Lin et al., 2003;
 692 Fasoli et al., 2018). Accordingly, an effective footprint-weighted mean flux estimate inferred from the receptor
 693 enhancement can differ from the unweighted afternoon-mean inventory flux over the sensitivity region. This
 694 difference depends on the spatial alignment between heterogeneous transport footprints and spatially heterogeneous

emissions (including localized hotspots), because under a given transport regime many grid cells within the domain may carry negligible footprint influence (Hüser et al., 2017; Kunik et al., 2019). We therefore interpret inventory comparisons as a plausibility envelope that reflects inter-inventory spread, rather than a validation target. Such inventory–observation differences are also reported for other coastal urban basins (e.g., Los Angeles) and are often sensitive to boundary-layer representation and meteorological inputs (Kim et al., 2025).

700

Beyond the winter fossil-fuel benchmark above, we further place the summer biogenic component and its offset ratio in the context of independent regional and urban estimates. Summer afternoons exhibited mean CO_2bio of $-3.59 \pm 0.63 \mu\text{mol m}^{-2} \text{s}^{-1}$, consistent with observation-based Pearl River Delta NEE (-0.1 to $-12 \mu\text{mol m}^{-2} \text{s}^{-1}$) (Mai et al., 2024a), and with modeled urban biogenic flux ranges (0 to $-15 \mu\text{mol m}^{-2} \text{s}^{-1}$) reported in previous studies (Wu et al., 2021; Wei et al., 2022; Kim et al., 2025). Consequently, summer-afternoon biogenic uptake offsets 60.13 % of concurrent CO_2ff at PY, with a bootstrap 95 % confidence interval of 48–72 %, highlighting substantial biogenic modulation of coastal urban CO_2 signals. Importantly, the inferred summertime offset remains substantial within the estimated uncertainty range, indicating robust biogenic modulation in magnitude. Comparable growing-season offsets have been reported for other coastal urban regions using independent approaches. A sensor-network combined with box-model analysis for Los Angeles suggests up to 60 % daytime offset; an inversion for the Boston coastal region indicates >50 % summer-afternoon offset (Kim et al., 2025; Sargent et al., 2018). High-resolution vegetation modeling for New York City similarly suggests ~40 % offset of afternoon anthropogenic enhancements and the potential to fully balance on-road traffic contributions (Wei et al., 2022). Overall, these contextual comparisons provide an external plausibility check and indicate that strong growing-season biogenic uptake is a plausible and important modulator of coastal-urban CO_2 signals in Guangzhou.

716 4 Conclusions

Three key knowledge gaps still limit the interpretation of atmospheric CO_2 signals in coastal megacities and their use for mitigation-relevant assessment. To address them, we present an observation-driven framework—without assimilating emission inventories—that combines multi-site CO_2/CO measurements, footprint-informed transport analysis, and a site-specific $\Delta\text{CO}/\Delta\text{CO}_2$ (R_{CO}) relationship to interpret coastal megacity CO_2 dynamics in Guangzhou (January 2023–September 2024). Relative to prior studies, this framework provides a distinct, process-based view of spatiotemporal CO_2 variability, sea–land breeze (SLB) modulation, and source–sink partitioning in a coastal setting. The three-site gradient reveals contrasting dominant controls by setting: transport governs the largest seasonal amplitude at the coastal site, biogenic exchange drives pronounced summertime daytime

drawdown at the vegetated site, and combustion dominates variability in the urban core. This combustion signal is characterized by a regression-derived R_{CO} , which is consistent with the broad post-2013 shift toward cleaner fuels and stringent vehicle emission standards in the Pearl River Delta. Importantly, these patterns point to a seasonally displaced coastal CO_2 “dome”. In contrast to the traditional paradigm where the maximum CO_2 enhancement is anchored over the urban center, our results demonstrate that in coastal megacities the dome’s peak can be displaced from the urban core by the combined effects of seasonal coastal transport/mixing and seasonally varying biogenic uptake associated with urban greening, underscoring the dynamic nature of coastal greenhouse gas distributions.

Under prevailing fair-weather coastal mesoscale conditions, the SLB circulation exerts a key control on diurnal variability in near-surface CO_2 and CO , with a non-linear seasonal modulation. In spring and winter, SLB strengthens ventilation and lowers CO_2 , whereas in summer it can favor accumulation (+2.08 ppm during sea-breeze hours) under weak-wind, stable conditions. Consistent with this, CO exhibits a more pronounced daytime enhancement than CO_2 during summer SLB days, supporting a trapping/recirculation regime in which combustion plumes are retained when coastal mesoscale flows coincide with shallow mixing and stable stratification. These seasonally opposing SLB impacts may be overlooked because many urban CO_2 studies focus on inland settings or annual-mean signals. Our results show that coastal mesoscale circulations can reverse the sign of SLB effects across seasons, highlighting important implications for inversion design and interpretation in coastal cities.

Source–sink attribution results at PY indicate that winter-afternoon CO_{2ff} in the urban core is robust within the quantified measurement and background-selection uncertainty ($\sim 3\%$ of the mean CO_{2ff}) and shows only small sensitivity to transport-model configurations ($\leq 1.47\%$ across the tested STILT setups), supporting stable winter-afternoon attribution under our sampling and background definition. In summer afternoons, the inferred CO_{2bio} ($-3.59 \pm 0.63 \mu\text{mol m}^{-2} \text{s}^{-1}$) indicates substantial biogenic uptake that offsets a large fraction ($\sim 60\%$) of the concurrent fossil contribution during the peak growing season, and this offset remains substantial within the estimated uncertainty bounds. The inferred summertime offset is broadly consistent with independent estimates reported for other coastal urban regions, providing an external plausibility check on the inferred source–sink separation.

Several limitations remain. The three-site network cannot resolve hyperlocal source heterogeneity, and SLB identification relies on near-surface wind criteria rather than full 3-D boundary-layer structure. Although configuration sensitivity is small, transport uncertainties associated with winds and boundary-layer mixing are not

756 fully quantified here. In addition, CO-based attribution is sensitive to variability in R_{CO} arising from changing
757 source mix, plume processing, and background definition, which propagates into CO_2^{ff} and the residual CO_2^{bio} .
758 Future work combining denser low-cost networks, boundary-layer profiling, and periodic isotopic constraints
759 would further tighten coastal urban carbon budgets.

760

761 Overall, our results demonstrate that coastal mesoscale dynamics can reshape both the magnitude and interpretation
762 of urban CO_2 signals, with SLB acting as ventilation in cool seasons but as a trapping/recirculation mechanism in
763 summer under weak-wind stability. The substantial summertime biogenic offset further indicates that biogenic
764 exchange associated with urban greening can materially modulate apparent fossil-fuel signals and should be
765 accounted for when interpreting mitigation trends. These findings have direct implications for coastal urban
766 monitoring and policy evaluation: to avoid season-dependent biases in assessing mitigation progress, urban
767 monitoring networks should prioritize the decoupling of biogenic signals—particularly during summer—to
768 accurately isolate anthropogenic contributions and thus ensure the robust evaluation of urban mitigation progress.
769 Furthermore, the strategic selection of monitoring sites in coastal megacities must explicitly account for the non-
770 linear accumulation effects of SLB circulations to ensure long-term sampling representativeness. While our
771 framework does not produce posterior flux fields as in a formal Bayesian inversion, it provides a complementary,
772 observation-driven tool for rapid process attribution and consistency checking in coastal urban carbon monitoring
773 and mitigation assessment.

774 **Code and data availability.** The STILT model source code used in this paper has been published on Zenodo and
775 can be accessed at <https://doi.org/10.5281/zenodo.1196561> (Fasoli, 2018). The EDGAR data used in this study are
776 publicly available at https://edgar.jrc.ec.europa.eu/dataset_ghg2024#conditions (last access: 18 June 2025)(Crippa
777 et al., 2024). The planetary boundary layer height data used in this study are available at
778 <https://doi.org/10.24381/cds.adbb2d47> (Hersbach, 2023). The NDVI data used in this study are available at
779 <https://doi.org/10.5067/MODIS/MOD13A3.006> (Didan, 2015). The CarbonTracker (CT-NRT.v2024-5) products
780 are available online at <https://doi.org/10.15138/ATPD-K925> (Jacobson et al., 2024). The NOAA Earth System
781 Research Laboratory/Global Monitoring Laboratory (NOAA GML) data used in this study are available at
782 <https://doi.org/10.25925/20241101> (Schuldt et al., 2024). Additional data and information used in this study are
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784 **Author contributions.** JWZ and ML designed the study. JWZ, YJL, CLP, BH, YYH, XFL, SJS, CLC, CW, ZZ,
785 JJJ and ML contributed to data collection and data analysis. JWZ designed and performed the model simulations.
786 JWZ and ML wrote the paper with contributions from all coauthors. JJJ and SJS provided valuable feedback and
787 opinions for paper refinement. All the authors revised the paper and edited the text.

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

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