Spatiotemporal patterns of the fossil-fuel CO2 signal in central Europe: results from a high-resolution atmospheric transport model

Journal Article

Author(s):
Liu, Yu; Gruber, Nicolas; Brunner, Dominik

Publication date:
2017-11-28

Permanent link:
https://doi.org/10.3929/ethz-b-000213663

Rights / license:
Creative Commons Attribution 3.0 Unported

Originally published in:
Atmospheric Chemistry and Physics 17, https://doi.org/10.5194/acp-17-14145-2017
Spatiotemporal patterns of the fossil-fuel CO$_2$ signal in central Europe: results from a high-resolution atmospheric transport model

Yu Liu$^{1,2}$, Nicolas Gruber$^{1,2}$, and Dominik Brunner$^3$

$^1$Environmental Physics, Institute of Biogeochemistry and Pollutant Dynamics, ETH Zurich, Zurich, Switzerland
$^2$Center for Climate Systems Modeling (C2SM), ETH Zurich, Zurich, Switzerland
$^3$Laboratory for Air Pollution/Environmental Technology, Swiss Federal Laboratories for Materials Science and Technology, Empa, Duebendorf, Switzerland

Correspondence to: Nicolas Gruber (nicolas.gruber@env.ethz.ch)

Received: 9 January 2017 – Discussion started: 13 April 2017
Revised: 29 September 2017 – Accepted: 16 October 2017 – Published: 28 November 2017

Abstract. The emission of CO$_2$ from the burning of fossil fuel is a prime determinant of variations in atmospheric CO$_2$. Here, we simulate this fossil-fuel signal together with the natural and background components with a regional high-resolution atmospheric transport model for central and southern Europe considering separately the emissions from different sectors and countries on the basis of emission inventories and hourly emission time functions. The simulated variations in atmospheric CO$_2$ agree very well with observation-based estimates, although the observed variance is slightly underestimated, particularly for the fossil-fuel component. Despite relatively rapid atmospheric mixing, the simulated fossil-fuel signal reveals distinct annual mean structures deep into the troposphere, reflecting the spatially dense aggregation of most emissions. The fossil-fuel signal accounts for more than half of the total (fossil fuel + biospheric + background) temporal variations in atmospheric CO$_2$ in most areas of northern and western central Europe, with the largest variations occurring on diurnal timescales owing to the combination of diurnal variations in emissions and atmospheric mixing and transport out of the surface layer. The covariance of the fossil-fuel emissions and atmospheric transport on diurnal timescales leads to a diurnal fossil-fuel rectifier effect of up to 9 ppm compared to a case with time-constant emissions. The spatial pattern of CO$_2$ from the different sectors largely reflects the distribution and relative magnitude of the corresponding emissions, with power plant emissions leaving the most distinguished mark. An exception is southern and western Europe, where the emissions from the transportation sector dominate the fossil-fuel signal. Most of the fossil-fuel CO$_2$ remains within the country responsible for the emission, although in smaller countries up to 80% of the fossil-fuel signal can come from abroad. A fossil-fuel emission reduction of 30% is clearly detectable for a surface-based observing system for atmospheric CO$_2$, while it is beyond the edge of detectability for the current generation of satellites with the exception of a few hotspot sites. Changes in variability in atmospheric CO$_2$ might open an additional door for the monitoring and verification of changes in fossil-fuel emissions, primarily for surface-based systems.

1 Introduction

With annual CO$_2$ emissions from fossil-fuel burning and cement production having soared in recent decades and approaching 10 Pg C yr$^{-1}$ (Raupach et al., 2007; Friedlingstein et al., 2014; Le Quéré et al., 2016), these fluxes have reached the same order of magnitude as the natural exchange fluxes between the atmosphere and land surface and between the atmosphere and the ocean, respectively (Sarmiento and Gruber, 2002; Le Quéré et al., 2016). Thus, the fossil-fuel emissions have become a key driver for the spatiotemporal dynamics of atmospheric CO$_2$, not only close to major sites of emissions but also far downstream (Peylin et al., 2011; Keppel-Aleks et al., 2013; Nassar et al., 2013). This represents simultaneously a challenge and an opportunity. It is an opportunity since the substantial and growing size of this fossil-fuel CO$_2$ signal facilitates the use of variations in atmospheric CO$_2$ to monitor and verify changes in fossil-fuel emissions.
two to three times larger than the differences resulting from intercomparison study, where inter-model differences in the advantage is well illustrated by the results of a recent model data but also on that of the transport model. The latter depends not only on the quality of the fossil-fuel emissions resulting accuracy of the modeled fossil-fuel CO\textsubscript{2} but this comes at the disadvantage that the processes underlying these emissions, ranging from the point-source nature of the emissions from coal-fired power plants, whose emissions vary in response to changing needs for electricity, to the strong diurnal fluctuations of the dispersed emissions associated with road transportation (Nassar et al., 2013). This strong spatial and temporal patterning of the fossil-fuel emissions interacts with the spatiotemporal variability of atmospheric transport, forming distinct patterns of the fossil-fuel signal in atmospheric CO\textsubscript{2} (Feng et al., 2016). Of particular relevance are the diurnal and the seasonal changes in emissions, since they tend to covary with atmospheric transport, which can lead to annual mean atmospheric CO\textsubscript{2} concentration gradients that differ from those attained if the emissions were held constant. This difference, which arises solely from the covariance between fluxes and transport, is called a “rectification effect” analogous to the rectification of an AC voltage in an electrical circuit by a diode (Denning et al., 1995; Zhang et al., 2016). Such unaccounted-for variations in the fossil-fuel signal would bias the biospheric signal in atmospheric inversion frameworks, hindering us from developing a better understanding of the role of the land biosphere as a carbon sink. At the same time, this strong temporal patterning of the emissions also creates distinct signals that might be used to detect or track the fossil-fuel signal.

In fact, several studies already explored the possibilities to detect the fossil-fuel signal and the emissions driving them (Ciais et al., 2014; Nassar et al., 2013). These include a range of methods and systems, including bottom-up methods based on surface observation systems (Shiga et al., 2014; McKain et al., 2012; Keller et al., 2016), CO and radiocarbon-based methods (Levin and Karstens, 2007; Van Der Laan et al., 2010; Vogel et al., 2013), airborne measurements (Turnbull et al., 2011), satellite constraints (Kort et al., 2012) and top-down approaches on the basis of atmospheric inversions (Ogle et al., 2015; Lauvaux et al., 2016; Brioude et al., 2013). Spatially, the focus ranges from point-scale (Bovensmann et al., 2010; Velazco et al., 2011; Turnbull et al., 2016) or urban-scale (Newman et al., 2013; Bréon et al., 2015; Turnbull et al., 2015; Pillai et al., 2016) to regional and global emissions (Keppel-Aleks et al., 2013; Basu et al., 2016).
exceeding a few 100 km. A successful detection also requires a good understanding of the contribution of the other processes affecting atmospheric CO$_2$ variations, namely the exchange fluxes with the land biosphere and with the ocean, respectively. Further, often it would be quite useful to know the source processes responsible for the fossil-fuel CO$_2$ signature, i.e., what fraction of the signal stems from emissions from a coal-fired power plant and what part from road transportation. This helps, for example, with the assessment of how the implementation of a particular policy affects the fossil-fuel signature, such as the shutting down of coal-fired power plants.

Few studies have taken a continental to global perspective on the fossil-fuel signal (Keppel-Aleks et al., 2013), as the focus in the last few years has been on urban areas (McKain et al., 2012; Newman et al., 2013; Kort et al., 2012), or just whether the emissions in the city can be detected or not (Hase et al., 2015; Pillai et al., 2016). In addition, comparatively less work has been carried out in Europe (Schneising et al., 2008), and the majority of it used relatively coarse-resolution atmospheric transport models, resulting in relatively washed-out gradients of the fossil-fuel signal over Europe (Keppel-Aleks et al., 2013; Peylin et al., 2011); a few of them focused on whether the potentially reduced emissions could be discerned by current observation methods in this region (Levin et al., 2011). Furthermore, little consideration has been given to the temporal variations of the emissions.

The main objective of this work is to fill these gaps and to develop a quantitative understanding of the fossil-fuel CO$_2$ signal in Europe. To this end, we employ a forward modeling approach using a high-resolution atmospheric transport model for Europe, forced with finely resolved fossil-fuel emission fluxes in time and space. In this paper, we will (i) investigate the magnitude of the contribution of the fossil-fuel CO$_2$ signal to the variations in total CO$_2$; (ii) understand how the high temporal resolution considered in the fossil-fuel emissions affect the fossil CO$_2$ signal; and (iii) determine the detectability of a reduction of fossil-fuel emissions from different sources through changes in the column mean CO$_2$ as seen, for example, by a satellite-based observing system. We first describe the model and methods, followed by the evaluation of the model. We then present the results, followed by a discussion of each of the aforementioned three main topics, and then conclude with a summary and an outlook.

2 Methods and data

To simulate the fossil-fuel CO$_2$ over central and southern Europe in the context of the variations in total atmospheric CO$_2$, we employ a regional high-resolution atmospheric transport model for the European domain and prescribe lateral and surface boundary conditions for the various components that constitute atmospheric CO$_2$. These include the fossil-fuel emissions, the CO$_2$ exchange fluxes with the land and ocean surfaces and the lateral atmospheric CO$_2$ boundary conditions. The simulations cover the period 27 March 2008 until 26 March 2009. The following subsections describe the methods and data in more detail.

2.1 Atmospheric transport model

The simulations were undertaken with the limited-area atmospheric prediction model COSMO (Consortium for Small-scale Modeling) (Baldauf et al., 2011) version 4.23. We employed the COSMO-7 setup developed by the Swiss Federal Office for Meteorology and Climatology (MeteoSwiss) for the purpose of providing boundary conditions for the inner COSMO-2 grid used for forecasting the weather in Switzerland. The COSMO-7 setup has a grid spacing of 6.6 km and its domain covers central and southern Europe (35.16$^\circ$N, 9.80$^\circ$E (lower left) to 56.84$^\circ$N, 23.02$^\circ$E (upper right); see Fig. 1).

The COSMO model is based on the primitive hydrothermodynamical equations describing compressible nonhydrostatic flow in a moist atmosphere without any scale approximations. The model equations are solved numerically on a rotated latitude–longitude grid, with terrain-following coordinates in the vertical (60 vertical levels, and lowest level at 10 m), using an Eulerian finite difference method. Parameterization schemes are used to resolve the sub-grid-scale physical processes such as vertical diffusion (turbulence), convection, radiation and soil processes. A tracer transport module was recently added to the COSMO model, permitting the online transport of passive tracers in a manner that is fully consistent with the physics of the model (Roches and Fuhrer, 2012). In our setup, advective transport was accomplished with a three-dimensional semi-Lagrangian scheme. The tracers are transported in the model as moist air mass mixing ratios $q_{CO_2}$. Values reported here are provided as dry air mole fractions $X_{CO_2}$, calculated as $X_{CO_2} = q_{CO_2}/(1-q_{H_2O}) M_{dry}/M_{CO_2}$, where $q_{H_2O}$ is the specific humidity and $M_{dry}$ and $M_{CO_2}$ are the molar masses of dry air and CO$_2$, respectively. The column-averaged dry air mixing ratio is calculated as follows:

$$X_{CO_2} = \frac{\sum_{k=1}^{K} (p(k + 1/2) - p(k - 1/2)) q_{CO_2}(k)}{\sum_{k=1}^{K} (p(k + 1/2) - p(k - 1/2)) (1 - q_{H_2O}(k))} \cdot \frac{M_{dry}}{M_{CO_2}},$$

where $k$ is the total number of vertical model levels ($K = 60$) and $p$ is pressure. We refer to Baldauf et al. (2011) for more details.
2.2 Fossil-fuel emissions

The fossil-fuel emissions for CO$_2$ were generated by merging a relatively coarse emission inventory for the regions outside Switzerland (EDGAR v4.2_FT2010, approx. 10 km; Janssens-Maenhout et al., 2012) with a high-resolution (0.5 km) emission inventory for Switzerland. The latter was produced by the company MeteoTest specifically for the CarboCount CH project. The annual emissions from this merged product for the year 2008 amount to 2.54 Pg CO$_2$ over the domain, representing about 10% of the global emissions of that year (Le Quéré et al., 2016). We merged the emission categories from the two inventories to five large emission categories: power generation, residential heating, road transportation, industrial processes, and others. Even though each of these different categories have a distinct emission pattern, many of them co-occur in the large metropolitan areas, leading to a very patchy emission pattern with strong emission hotspots and extensive regions with relatively low emission densities (Fig. 1).

These emission inventories are given for each emission category as annual totals for each grid cell, i.e., $E_{\text{ann}}$, requiring us to multiply them with time functions to generate hourly time series of the fossil-fuel emissions at each location (Nassar et al., 2013). The time functions we employed were originally generated by the University of Stuttgart (Institut für Energiewirtschaft und Rationelle Energieanwendung, IER) for the GENEMIS project (Friedrich and Reis, 2004) and have been used since in several air quality modeling studies. The time functions are comprised of diurnal, weekly and seasonal components and are specific to each of the main economic sectors (activities collected in the Selected Nomenclature for Air Pollution (SNAP) codes) (Kuenen et al., 2014). When constructing these time functions, it is ensured that their annual mean is equal to unity so that the annual totals remain unchanged. The scaled emission flux, $E(t)$, is then computed by modifying the annual total fossil-fuel emission, i.e., $E_{\text{ann}}$:

$$E(t) = E_{\text{ann}} \cdot f_{\text{diurnal}}(t) \cdot f_{\text{week}}(t) \cdot f_{\text{season}}(t),$$

(2)

where $t$ is time (hour of the year) and $f_{\text{diurnal}}$, $f_{\text{week}}$ and $f_{\text{season}}$ are the diurnal, weekly and seasonal scaling factors, respectively. The time function factor $f_{\text{diurnal}}$ depends on the hour of the day (local time (LT), $t_{\text{hour}} = t$ modulo 24 h) and is different for weekdays and weekends to reflect the different level of activities on weekdays and weekends. The factor $f_{\text{week}}$ depends on the day of the week, with one value for weekdays (Monday–Friday) and a lower one for Saturdays and Sundays. The factor $f_{\text{season}}$ depends on the month, but in order to avoid discontinuities between subsequent months it is linearly interpolated to a given day between the centers (day 15) of 2 adjacent months. The time functions (except for the daily one) vary also by country and are locally adjusted to reflect local time. Some reassignments were nec-
Figure 2. Time dependence of fossil-fuel CO\textsubscript{2} emissions for different sectors and countries. (a) Time functions for the diurnal and weekly emissions for four sectors. (b) Annual evolution of the CO\textsubscript{2} emission intensity for Germany, Switzerland and southwest Europe (Portugal and Spain). Panel (c) is the same as panel (b) but for the domain total. Panel (d) is as panel (b) but for United Kingdom, Italy and eastern Europe (Poland, Czech Republic, Slovakia and Hungary). The range shown in panels (b)–(d) are the daily minima and maxima for each country or group of countries.

necessary to align the categories used in EDGAR v4.2 and the CarboCount CH inventory (both following IPCC guidelines) with the SNAP categories as described in the Supplement.

The time functions differ greatly between the various categories, reflecting their very different time course of activities over the average day, week or year (see Fig. 2a and b). Among all diurnal time functions, road transportation has the largest diurnal variability and is characterized by two peaks during the day that reflect the rush hour periods (08:00–09:00 and 17:00–18:00 LT). Also residential and commercial combustion have a distinct diurnal cycle with two peaks. In contrast, the emissions from industrial processes and fossil-fuel-fired power plants vary less over the course of the day and also have only one peak. The time functions for the day-of-week reflect primarily the lower industrial activities and traffic during the weekend, while most other sectors continue to emit at only slightly smaller rates (see Fig. 2a). Combining all the sectors together, emissions during the weekend are 15–20\% lower than during the week. The seasonal time functions depend primarily on the local climatic conditions (see Fig. 2b and d), with northern, eastern and central European countries having a maximum in winter, likely due to their need for residential heating, while there is little seasonality in emissions in the southern European countries. The seasonality of the total emission (Fig. 2c) follows that of the northern, eastern and central European countries with a wintertime maximum, although somewhat less pronounced due to leveraging effect from the southern European countries.

In order to be able to trace the fossil-fuel signature in atmospheric CO\textsubscript{2} back to the emitters, we consider separate fossil-fuel tracers for 10 different countries (or groups of countries) in our atmospheric transport model (see Fig. 1). Each of these tracers receives only the emissions from its respective country or group of countries, while elsewhere the emissions are set to zero. Due to the linearity of atmospheric transport and the absence of any transformation of CO\textsubscript{2} in the atmosphere, the individual country-based tracers can then be summed to obtain the total fossil-fuel CO\textsubscript{2} signal. In addition, in order to determine the contribution of the different CO\textsubscript{2} emission categories to the total fossil-fuel CO\textsubscript{2}, we also included five additional fossil-fuel tracers, one each for the five categories we consider: power generation, residential heating, road transportation, industrial processes, and others. For these five tracers, we used time-invariant emissions, permitting us to assess also the role of the time variations in emissions on the fossil-fuel CO\textsubscript{2} signal. In total, we included
17 fossil-fuel tracers (10 countries, 5 sectors, total fossil-fuel CO\textsubscript{2} with time-varying emission and total fossil-fuel CO\textsubscript{2} with time-constant emission) in our high-resolution simulation study.

At the lateral boundaries, we employ a partial relaxation boundary condition for these 17 tracers. In such a partial relaxation, the tracer is relaxed to the boundary concentration only at the outermost grid cells of the domain and only when the wind is directed toward the inside of the domain (in COSMO, this option is provided by the switch “T_RELAX_INFLOW”). Since we are interested in the fossil-fuel signal emanating from emissions in Europe only, the lateral boundary concentration was set to zero. Through this option, we avoid creating a situation where the zero concentration boundary condition is propagated (erroneously) against the flow back into our domain.

2.3 Other CO\textsubscript{2} component fluxes

In order to simulate the distribution of total atmospheric CO\textsubscript{2}, we also include in our model three other CO\textsubscript{2} components, namely background CO\textsubscript{2}, the terrestrial biospheric CO\textsubscript{2} and the oceanic CO\textsubscript{2} components. The background CO\textsubscript{2} represents that part of the atmospheric CO\textsubscript{2} that enters the domain through its boundaries. These boundary concentrations are provided by the post-assimilation results of CarbonTracker Europe (Peters et al., 2010). For this tracer, we use a “full” relaxation boundary condition. This means that we are restoring the modeled mixing ratio toward the value provided by CarbonTracker across a transition zone consisting of 13 grid cells, with the relaxation increasing in strength from the inner to the outer border of this zone. In COSMO, this option is provided by the “T_RELAX_FULL” switch.

For the terrestrial biospheric CO\textsubscript{2} component, we used the hourly terrestrial biospheric fluxes from the Vegetation Photosynthesis and Respiration Model (VPRM) (Mahadevan et al., 2008). For the oceanic CO\textsubscript{2} component, we combined the monthly air–sea CO\textsubscript{2} flux estimates for the Atlantic from Landschützer et al. (2013) with the annual mean flux estimates for the Mediterranean by D’Ortenzio et al. (2008). As the oceanic flux contribution is small, no attempt was made to add higher-frequency variability. The lateral boundary conditions for these two tracers were handled the same way as those for the fossil-fuel signal, i.e., a partial relaxation toward a zero concentration at the boundary.

2.4 Simulations

The hindcast simulation started on 1 March 2008, with the initial and boundary conditions for meteorology taken from the operational hourly COSMO-7 analyses of MeteoSwiss and the initial and boundary conditions for atmospheric CO\textsubscript{2} provided by CarbonTracker Europe (Peters et al., 2010). The model was then run for 13 months until 26 April 2009. No assimilation of any meteorological data was performed.

The lateral and surface boundary conditions for the total of 21 CO\textsubscript{2} tracers considered (17 fossil fuel, 4 other components) were prescribed as described above. We consider the first 26 days as a spinup and use the subsequent 12 months for our analyses.

3 Evaluation

3.1 Total atmospheric CO\textsubscript{2}

We evaluate our COSMO-based results for the total atmospheric CO\textsubscript{2} concentration (computed by summing the fossil-fuel component with the three others) by comparing them to the measurements from four sites in central Europe, namely Mace Head (MHD; 33.33°N, 9.90°W; 5 m above ground, 15 m a.s.l.; coastal site), Cabauw (CBW; 51.97°N, 4.92°E; 20, 60, 200 m above ground, 0 m a.s.l.; flatland, near urban site), Hegyhatsal (HUN; 6.95°N, 16.65°E; 10, 48, and 115 m above ground, 248 m a.s.l.; continental site) (Geels et al., 2007) and Puy de Dôme (PUY; 45.46°N, 2.58°E, 1480 m a.s.l.; mountain site).

The modeled daily mean atmospheric CO\textsubscript{2} at these four sites agrees generally well with the corresponding observations, although the agreement deteriorates with proximity to the ground (see Table 1). This deterioration is likely a consequence of the stronger influence of local processes closer to the ground, which are more difficult to capture by the model. At the highest measurement level, the correlation between the modeled and observed values exceed 0.6 at all sites and are statistically significant at the $p < 0.05$ level. The highest correlation is found at MHD ($r = 0.80$). This is due to the relatively steady conditions that characterize this relatively clean coastal site. Influence from air pollution is only observed during episodes of transport from the UK and continental Europe, which are very well captured by the model. The correlations are somewhat lower at the more polluted or more continental sites, i.e., between 0.57 (lowest level) and 0.63 (highest level) at the coastal tall tower station CBW in the Netherlands and between 0.52 (lowest level) and 0.68 (highest level) at the continental tall tower station HUN in Hungary. The atmospheric CO\textsubscript{2} variations at the mountain top site PUY in France are well captured ($r = 0.72$). COSMO tends to systematically underestimate the observed CO\textsubscript{2} concentration at most of the stations and levels, except at the coast of Ireland (MHD), where it is overestimated by 0.3 ppm (Table 1).

The biases tend to get larger with increasing continentality of the sites and the associated higher complexity of the surrounding terrain and other influencing factors. At CBW, the biases amount to between $-1.4$ and $-5.7$ ppm, while in central Hungary (HUN) the biases exceed $-4$ ppm at even the highest vertical level.

The general underestimation of the total CO\textsubscript{2} can stem from biases in any of the components that make up the total CO\textsubscript{2} plus biases in atmospheric transport and mixing. The
Table 1. Evaluation of COSMO-7-based simulations of the atmospheric CO$_2$ concentration at four European sites. The comparison (observations minus model) are computed using the daily means for the period of 27 March 2008 through 26 March 2009.

<table>
<thead>
<tr>
<th>Station</th>
<th>Characteristics</th>
<th>Relative height (m)</th>
<th>SD obs (ppm)</th>
<th>SD mod (ppm)</th>
<th>Correlation</th>
<th>Bias (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cabauw (CBW, Netherlands)</td>
<td>tower</td>
<td>20</td>
<td>17.11</td>
<td>12.05</td>
<td>0.57</td>
<td>-5.66</td>
</tr>
<tr>
<td>Cabauw (CBW, Netherlands)</td>
<td>tower</td>
<td>60</td>
<td>12.94</td>
<td>11.26</td>
<td>0.61</td>
<td>-2.32</td>
</tr>
<tr>
<td>Cabauw (CBW, Netherlands)</td>
<td>tower</td>
<td>200</td>
<td>9.82</td>
<td>7.95</td>
<td>0.63</td>
<td>-1.45</td>
</tr>
<tr>
<td>Puy de Dôme (PUY, France)</td>
<td>mountain top</td>
<td>10</td>
<td>7.14</td>
<td>6.82</td>
<td>0.72</td>
<td>-0.8</td>
</tr>
<tr>
<td>Hegyhatsal (HUN, Hungary)</td>
<td>continental</td>
<td>10</td>
<td>19.18</td>
<td>9.02</td>
<td>0.52</td>
<td>-12.37</td>
</tr>
<tr>
<td>Hegyhatsal (HUN, Hungary)</td>
<td>continental</td>
<td>48</td>
<td>12.99</td>
<td>8.78</td>
<td>0.6</td>
<td>-7.31</td>
</tr>
<tr>
<td>Hegyhatsal (HUN, Hungary)</td>
<td>continental</td>
<td>115</td>
<td>10.54</td>
<td>8.09</td>
<td>0.68</td>
<td>-4.14</td>
</tr>
<tr>
<td>Mace Head (MHD, Ireland)</td>
<td>coastal</td>
<td>15</td>
<td>6.8</td>
<td>3.92</td>
<td>0.80</td>
<td>0.33</td>
</tr>
</tbody>
</table>

low and positive bias at MHD, where the contribution of the background CO$_2$ component dominates, suggests that this component is overall well modeled and likely not responsible for the bias at the other sites. Since the contribution of the oceanic fluxes is very small, this component can be excluded as an explanation as well. Thus, the general underestimation is thus likely due to the superposition of biases in atmospheric transport and biases in the underlying boundary conditions for the fossil-fuel emissions and/or terrestrial fluxes. COSMO is known for ventilating the planetary boundary layer (PBL) too strongly, particularly in wintertime under weakly stratified conditions. This may be especially acute for the HUN site, because the air in the lowest atmospheric levels tends to get trapped at this site owing to the wintertime prevalence of anticyclonic conditions in the Carpathian Basin (Haszpra et al., 2012). An alternative explanation is that the biospheric sink simulated by VPRM is too strong, as discussed later. Unfortunately, we do not have observationally based estimates of the fossil-fuel or terrestrial biosphere components at the four sites discussed so far, requiring us to use data from other sites for further evaluation.

Even though COSMO exhibits some biases in the mean, it captures the observed variability generally well (Table 1). In particular, COSMO reproduces the strong gradient in variability between the coastal site MHD (obs: ~7 ppm; mod: ~4 ppm) and the continental site in central Hungary (obs: ~11 ppm; mod: ~8 ppm), reflecting primarily differing contributions of synoptic variations on atmospheric CO$_2$. However, the absolute magnitude of the variability is not well captured by our simulations, with COSMO consistently underestimating the observed variability. Overall, the evaluation of the total atmospheric CO$_2$ concentration reveals an agreement with the observations at the four sites in terms of both mean and variability. The agreement is clearly better further away from the ground, i.e., at heights of at least 50 m above the ground.

3.2 Fossil-fuel CO$_2$ component

Estimates of the fossil-fuel component in atmospheric CO$_2$ are available for our model simulation period from Lutjewad in the Netherlands (LUT; 6.35°E, 53.4°N; 1 m a.s.l.) (Van Der Laan et al., 2010; Bozhinova et al., 2014) and from Heidelberg in Germany (HEI; 49.417°N, 8.675°E; 116 m a.s.l.) (Levin and Karstens, 2007). Both estimates are based on a combination of concurrent CO and $^{14}$CO$_2$ measurements and represent the fossil-fuel-induced offset relative to a regional background. They are thus comparable to our modeled fossil-fuel component, as this reflects the offset relative to the domain-wide background induced by the lateral boundary conditions. Lutjewad is located on the Wadden Sea dike in the north of the Netherlands, influenced by the highly populated and industrialized areas in the Netherlands and in northwestern Germany (Ruhr area). The Heidelberg station is located near an urban center with considerable fossil-fuel emissions.

At the Dutch site LUT, the daily-averaged fossil-fuel CO$_2$ component simulated by our model compares well with the observations ($r = 0.73$, mean bias ~4 ppm) (see Fig. 3a). Generally, the model reproduces the observed variability, especially in summer, when the fossil-fuel CO$_2$ component is low due to the deep mixing in the atmosphere. But the model underestimates the estimated fossil-fuel CO$_2$ component substantially in winter. This may be due to several reasons. First, the model may transport signals too quickly out of the PBL, which is a known problem of many atmospheric transport models under stratified conditions typical of wintertime (see also above) (Holtslag et al., 2013). Second, our wintertime emission inventory in the region might be too small due to, for example, our underestimating the strength of the seasonal signal in the time functions. Third, the observations might be biased high. One reason is that these reconstructions rely on a constant ratio between CO and $^{14}$CO$_2$, which may lead to an underestimation of the $^{14}$C–CO ratio compared to real values at some time of the year and subse-
quent overestimation of the inferred fossil-fuel CO$_2$ (Van Der Laan et al., 2010; Bozhinova et al., 2014).

At Heidelberg, our model captures the fossil-fuel CO$_2$ component even better, particularly since the model has a very small mean bias of 0.75 ppm. Also the day-to-day and the seasonal variations are well represented with a correlation coefficient of 0.72. The model’s (small) overestimation of the fossil-fuel component may be due to our prescribing all emissions at the surface, while the fossil-fuel-fired power plants that contribute substantially to the fossil-fuel CO$_2$ at this site tend to have an effective emission height quite some distance above the ground due to the height of the stacks and the additional rise of the buoyant plumes (Vogel et al., 2013). Another reason might be an overestimation of the emissions in our emission inventory EDGAR – an explanation furthered by EDGAR’s emission being higher than those of IER (Peylin et al., 2011). Especially assuring, and particularly so in comparison to the situation at LUT, is the COSMO model’s ability at HEI to capture most of the variability and amplitude of the fossil-fuel component in winter. An exception are the observations from late December and early January, where the data include a number of exceptionally high peaks. These peaks may be the result of very strong local trapping of the emitted fossil-fuel CO$_2$ by, for example, a local inversion situation, i.e., a process that our model cannot properly resolve.

Despite these discrepancies, the evaluation results provide us with good confidence to use our COSMO-7-based system to investigate the spatiotemporal variability of the fossil-fuel CO$_2$ in central and southern Europe. The presence of an overall negative bias in the total atmospheric CO$_2$ in the absence of such a bias in the fossil-fuel component suggests that the bias comes from the terrestrial biospheric component. This could be due to our VPRM-based estimates of the net fluxes being too negative, as suggested by Oney et al. (2017), i.e., suggesting a too strong sink for central and southern Europe, or for our model simulating a too small diurnal and/or seasonal rectification effect (Denning et al., 1995), i.e., a too small correlation between the time variations in the terrestrial exchange fluxes and atmospheric transport and mixing. This deficiency does not impact our results much, since our focus will be on the spatiotemporal variability of the fossil-fuel CO$_2$ signal.

4 The spatiotemporal pattern of the fossil-fuel CO$_2$

4.1 The spatial pattern

In the annual mean, the fossil-fuel component of atmospheric CO$_2$ in the surface layer (∼10 m above ground) amounts to more than 10 ppm across wide swaths of central Europe (Fig. 4a). We computed this mean using data from all times of the day in order to fully reflect the annual mean. In large metropolitan areas, such as in western Germany (Ruhr region), Berlin, London, Paris and Milan, the annual mean fossil-fuel component exceeds even 30 ppm. To first order, the distribution of the surface fossil-fuel CO$_2$ reflects the distribution of the emissions (see Fig. 1), suggesting a somewhat limited effectiveness of atmospheric transport and mixing to disperse the signal aloft and in lateral directions. In mountainous regions this is clearly a consequence of topographic constraints; elsewhere this is largely a result of the strong spatial gradients in emissions, which remain conserved in the annual mean due to the overall diffusive nature of the dispersion. Nevertheless, a substantial amount of the emitted CO$_2$ is being transported away, leading to a sizeable fossil-

Figure 3. Comparison between modeled and observation-based estimates of the fossil-fuel CO$_2$ component for the years 2008–2009. (a) Comparison at the Lutjewad site in the Netherlands (LUT; 6°21′ E, 53°24′ N; 1 m a.s.l.) (Van Der Laan et al., 2010; Bozhinova et al., 2014). (b) Comparison at Heidelberg (HEI; 49.417° N, 8.675° E; 116 m a.s.l.) (Levin and Karstens, 2007). The observational estimates are based on concurrent observations of CO and $^{14}$CO$_2$. 

Atmos. Chem. Phys., 17, 14145–14169, 2017

www.atmos-chem-phys.net/17/14145/2017/
fuel CO$_2$ signal extending far into the oceans surrounding Europe, especially the North Sea.

Despite this lateral transport, the relatively good conservation of the spatial gradients in emissions sets our results distinctly apart from previous studies, where the fossil-fuel CO$_2$ signal was modeled to be very smooth in space and on average also substantially smaller. For example, compared to the results obtained with the medium-resolution (0.5°) Regional Model (REMO) (Peylin et al., 2011), one can detect in our simulations nearly all major metropolitan regions and other fine-scale features, such as individual fossil-fuel-fired power plants (e.g., in eastern Germany). This is primarily the result of the high horizontal and vertical resolution of COSMO permitting this model to conserve the spatial gradients well. This good conservation is particularly well illustrated when considering snapshot distributions of the fossil-fuel CO$_2$ for individual seasons (Fig. 5). This figure also shows the strong impact of the transport and dilution by the diurnal variations of the PBL, whose impact is particularly strong in summer.

For much of Europe, the fossil-fuel component is the dominant contributor to the spatial gradients in annual mean atmospheric CO$_2$ (Fig. 4b–d). In many places it accounts for nearly all of the spatial gradients, with the contribution of the background and the terrestrial biospheric component being substantially smaller. The latter shows gradients up to 10 ppm (Fig. 4c), while the background signal does not exceed a few ppm (Fig. 4d). In the big cities, the fossil-fuel CO$_2$ component represents even a sizeable fraction (10%) of the total CO$_2$ concentration. This dominance of the fossil-fuel component together with its highly patterned nature owing to the many point sources leads to a hotspot pattern in the near-surface map of total atmospheric CO$_2$ over much of Europe (Fig. 4b). However, due to lower emissions in southwestern Europe, the fossil-fuel CO$_2$ signal is less strikingly visible there compared to central Europe. At the same time, the sign of the biospheric signal changes in the south and becomes positive. This compensates for the smaller fossil-fuel signal there and results in a relatively uniform spatial pattern of atmospheric CO$_2$ across Europe (Fig. 4b). Also the relatively low CO$_2$ concentrations in the mountain regions, such as the Alps, Apennines, Pyrenees and central France, reflect the much lower contribution from the fossil-fuel component.

Naturally, when investigating the column-averaged dry air mole fractions ($X_{CO_2}$), i.e., the property typically measured...
by remote sensing from a satellite, the annual mean gradients of the fossil-fuel component are much smaller than those seen at the surface (see Fig. 6a). This is a consequence of the lateral gradients being much weaker aloft due to a more effective horizontal transport and mixing. An additional reason is a much stronger influence of the lateral boundary conditions, which results in a dilution of the fossil-fuel components. As a result, most of the hotspot nature seen in the surface concentration pattern is blurred in $XCO_2$. Also the magnitude of the signal is much weaker. While the surface signal of the fossil-fuel CO$_2$ signal amounted to more than 30 ppm in strong emissions regions, the signal in the column-averaged annual mean $XCO_2$ hardly exceeds 2 ppm. The impact of the predominant westerly air flow becomes much more obvious in the column-averaged dry air mole fraction $XCO_2$, with the fossil-fuel component revealing a clear eastward increase that is substantially stronger than the gradient in the underlying emissions.

The relative dominance of the fossil-fuel component over the other components of atmospheric CO$_2$ is much weaker when considering the column-averaged dry air mole fraction of CO$_2$ (see Fig. 6b–d). As a result, the total $XCO_2$ is made up of three relatively equally sized contributions, with the fossil-fuel CO$_2$ signal continuing to dominate the $XCO_2$ variations in the major metropolitan areas. Contrary to the annual surface pattern, where CO$_2$ tends to increase eastward, the highest $XCO_2$ are found in southwestern Europe with a trend toward lower values going eastward. This is partly a consequence of the lateral boundary conditions for atmospheric CO$_2$, which tend to lead to the advection of elevated background CO$_2$ into the domain from the southwest. But the most important reason is the strong negative terrestrial biosphere signal over Europe, reflecting the sizeable carbon sink in European forests in the last decade (Reuter et al., 2017). Interestingly, the relatively uniform negative distribution for $XCO_2$ in Fig. 6c contrasts with a more patterned biospheric signal in the lowest layer of the atmosphere (Fig. 4c). There, the strong negative signal is restricted to central Europe, while much of southern Europe has a positive annual mean biospheric signal. The likely reason for this difference is the biospheric rectification effect (Denning et al., 1995), which tends to lead a vertical redistribution of CO$_2$, i.e., positive values in the lower atmosphere and negative ones aloft. In most of Europe, this rectification signal is relatively small in comparison to the annual mean biospheric component, so that the latter determines the overall signal. But in southern Europe, where the biospheric fluxes tend to be smaller in magnitude and in the annual mean to be near zero, the
rectifier effect can dominate, explaining the positive signals in the surface layer (Fig. 4c) and simultaneously the negative signals when the biospheric signal is integrated vertically (Fig. 6c).

4.2 The temporal variability

The temporal variability of the fossil-fuel CO$_2$ signal at the surface is very large, leading to a standard deviation around the annual mean of 30 ppm or more in the hotspot regions (Fig. 7a). These hotspots correspond largely to the regions of highest emissions (Fig. 1). This high variability is a result not only of the temporal variability of the emissions but also of the interaction of variability in atmospheric transport and mixing with the strong lateral gradients seen in the snapshot figures (see Fig. 5).

A similar pattern of variability is seen in surface atmospheric CO$_2$ (Fig. 7b), suggesting that the fossil-fuel CO$_2$ is a major determinant not only of the annual mean spatial distribution of atmospheric CO$_2$ but also of its temporal variability. This is confirmed by Fig. 8a, which shows the relative contribution of the fossil-fuel CO$_2$ signal to the temporal standard deviations of atmospheric CO$_2$. In many places, particularly in Europe’s major metropolitan areas but also in many urban areas across Europe, the fossil-fuel signal dominates the variability in atmospheric CO$_2$. But the high fossil-fuel contribution is not limited to the urban areas. In fact, the region delineated by having a 50% contribution or more extends over much of northern central Europe, including the North Sea (see Fig. 8a).

In order to better understand the origin of the strong variability, we decomposed the variability into seasonal, synoptic and diurnal contributions. The seasonal variation component was derived by averaging the data on a monthly basis and by subtracting the annual mean. The synoptic component was then computed by subtracting from the data the time series of the monthly means and then forming daily averages of these deseasonalized data. Finally, the diurnal variability was derived by subtracting the seasonal and synoptic components from the data. To determine the fractional contribution, we then computed the fractional variance of each component relative to the total variance. Since the different temporal components can compensate for each other, the sum of the fractional variance can actually exceed unity.

This decomposition reveals that the contribution of the fossil-fuel CO$_2$ to the total variability of atmospheric CO$_2$ varies greatly depending on the temporal scale considered (Fig. 8). While the fossil-fuel contribution is comparably small on seasonal timescales (Fig. 8b), the contribution on synoptic and particularly on diurnal timescales is actually
Figure 7. Maps of the annual standard deviation of (a) the fossil-fuel component and (b) atmospheric CO$_2$ in the surface layer. Shown are the results for the period 27 March 2008 through 26 March 2009.

Figure 8. Maps of the contribution of fossil-fuel CO$_2$ variability to total atmospheric CO$_2$ variability within the lowest model layer (0–20 m, centered at 10 m) on various timescales in percent: (a) contribution on all timescales; (b) contribution for the seasonal timescale only; (c) contribution for the synoptic timescale only; (d) contribution for the diurnal timescale only. Note that the contributions from panels (b) through (d) do not add up to the numbers shown in panel (a). This is a result of a partial compensation between the different temporal components due to the temporal covariations in fossil-fuel and total atmospheric CO$_2$. 
very large, exceeding 60% across nearly the entire northern part of central Europe (Fig. 8c and d). The small contribution on the seasonal timescales is the result of the dominance of the seasonal cycle of the biospheric fluxes in most of Europe. An exception are a few places in northern Europe and in the very south of our European domain. We interpret this to be caused primarily by the relatively strong seasonality of the fossil-fuel emissions in these regions due to the strong summertime requirement for cooling in the south and the strong wintertime demand for heating in the north.

The pattern of the fossil-fuel contribution on synoptic timescales is very similar to that of the total contribution, meaning its contribution dominates the total temporal variability. This is consistent with synoptic variations also being among the strongest contributors to atmospheric variability due to baroclinic waves and frontal systems being formed out of the strong baroclinicity that characterize the mid-latitudes. These synoptic weather events transport the emitted CO$_2$ also quite efficiently outside the main metropolitan areas, explaining the widespread signal of the fossil-fuel contribution to the total variance of atmospheric CO$_2$. Even larger than the fossil contribution to synoptic variability is the contribution on the diurnal timescale, with the fossil-fuel CO$_2$ contributing more than half of the variability over most of Europe. This high variability comes from the interaction of the diurnal variability of the fossil-fuel emissions, with the strong diurnal variability of atmospheric transport, particularly the diurnal mixing of the PBL. This covariability between fossil-fuel emissions and atmospheric transport exceeds that between the biospheric fluxes and atmospheric transport over the entire year due to the latter fluxes being large and relevant only during the spring–summer period, while the fossil-fuel emissions are relatively high during most of the months of the year, particularly close to the sources.

5 Discussion

The analyses of the results raise a number of questions that we would like to discuss next. First, why is the diurnal variability so high and, in particular, what is the contribution of the diurnal (and seasonal) variations in CO$_2$ emissions on the simulated fossil-fuel CO$_2$ signal? Further, is there an impact beyond the variability, e.g., on the mean fossil-fuel CO$_2$ signal? Second, what is the contribution of the various sectors on the fossil-fuel CO$_2$ signal and in what way do emissions from one country influence the fossil-fuel CO$_2$ signal in another country? Third, how can we use the insights gained from the study of the fossil-fuel CO$_2$ signal to develop optimal strategies for detecting changes in fossil-fuel CO$_2$ emissions? We discuss each of these three questions next.

5.1 The impact of variations in fossil-fuel emissions on atmospheric CO$_2$

In order to elucidate the role of the temporal variations in fossil-fuel emissions on the fossil-fuel CO$_2$, we contrast the results of our standard simulation with time-varying emissions with those where the fossil-fuel emissions were kept constant over time. The annual emissions are identical for the two cases, but the time-constant case has, on average, considerably higher emissions in summer and at night. The contrast between these two cases shows only a small change in the high diurnal variability of atmospheric CO$_2$ seen in Fig. 8d (results not shown). The largest changes are found around some of the large metropolitan areas (e.g., London, Paris, Milan), but they do not exceed 10%. Thus the majority of the diurnal variability in the fossil-fuel CO$_2$ stems from the diurnal variations in atmospheric transport and mixing acting on the strong horizontal gradients in emissions.

While not contributing much to the diurnal variability in the fossil-fuel CO$_2$, the consideration of the time-varying emission matters quite substantially for the annual mean distribution of the fossil CO$_2$ signal. Figure 9a reveals that the annual mean fossil CO$_2$ signal in the simulation with time-varying emissions is substantially lower over wide swaths of Spain, Italy, Benelux, (western) Germany and the UK compared to the simulation where fossil-fuel emissions were kept constant. The strongest negative signals are found close to the strongest emitters in these countries, with magnitudes exceeding several ppm. But the magnitude of the signal does not correspond to the magnitude of emissions, since regions with comparably low emissions such as Spain have signals that are as large as those in high emission regions of the Netherlands. The relatively large signals in southern Europe are likely due to the stronger PBL dynamics in these regions throughout the year in comparison to central and northern Europe. Some regions also have a positive signal from the time-varying emissions, such as parts of France and northeastern Germany. Thus the interaction between the variations in fossil-fuel emissions and the variations in atmospheric transport and mixing leads to a substantial net signal in atmospheric CO$_2$, even though the total emissions in both cases are identical.

This net signal represents a fossil-fuel-driven rectification effect (Zhang et al., 2016) in analogy to the rectification effect associated with the terrestrial biosphere (Denning et al., 1996; Larson and Volkmer, 2008), i.e., a signal that is due to the covariance of emissions and atmospheric transport and mixing. Its (mostly) negative sign emerges from the fact that when the emissions are large, e.g., during the day, the transport and mixing away from the surface is strong, diluting the fossil-fuel signal in atmospheric CO$_2$. In contrast, when the emissions are small, e.g., during the night, the transport and mixing tends to be weak. Taken together, this results in a more efficient dilution of the emissions in the time-varying emission case compared to the time-invariant case, thus ex-
Figure 9. Maps of the impact of the consideration of time-varying fossil-fuel emissions. (a) Difference in annual mean surface CO$_2$ between the case with time-varying and time-constant fossil-fuel emissions. This difference represents the fossil-fuel rectification effect. (b) Linear correlation between the fossil-fuel emissions and the height of the planetary boundary layer height in the COSMO-7 model. Pixels with emissions smaller than 0.06 gC m$^{-2}$ yr$^{-1}$ are not plotted. The positive correlation implies high emissions when the PBL is deep, and vice versa. Most of this correlation stems from the diurnal timescale, but the correlation is enhanced through the (mostly) positive correlation also on seasonal timescales (see main text). The negative correlations over the ocean stem from the fossil-fuel emissions by ships.

plaining the mostly negative sign of the fossil-fuel rectification effect.

This explanation is supported by the mostly positive correlation between the height of the PBL and the fossil-fuel emissions, since the height of the PBL is a good proxy for the magnitude of the mixing and transport in the lowest levels of the atmosphere (Fig. 9b).

But there are a number of notable exceptions. For example, wide swaths of northeastern Germany and Poland and some places in central France have a positive rectification signal. Further, there are places where the covariation of fossil-fuel emission and the PBL is negative, yet the fossil-fuel rectification effect is still negative (e.g., the Ruhr valley region in western Germany), suggesting that our explanation does not cover all aspects. In response, one first needs to recognize that not only PBL but also other temporally varying phenomena, such as local atmospheric circulation patterns (e.g., mountain winds, sea breezes), can lead to covariability between emissions and transport/mixing, creating a rectification signal that can differ in sign. The contribution of the sea breeze can be identified quite clearly by the strong negative sign along most of the coastline between southern Europe and the Mediterranean. Second, the local timing between the growth and decay of the PBL and the emissions can be quite different due, in part, to the substantially different time functions for the different emission categories and their different local contributions (Fig. 1). For example, in regions with a large contribution from road transportation, the local emissions have a strong peak in the early morning hours, when the PBL is still shallow, leading to a high signal there, while emissions are lower when the PBL is at its maximum in the early afternoon. This would create a positive rectification signal. Finally, in certain places, also the seasonal rectification appears to play a role, i.e., the seasonal covariations of the emissions with the PBL height. In fact, in many places the magnitude of the correlation between emission and PBL height on seasonal timescales exceeds that on diurnal timescales. This seasonal variation is particularly large for residential heating, which is maximum in winter when the PBL is low, leading to a positive seasonal rectification. This effect likely contributes to the negative correlations between emissions and PBL height in large urban centers such as Paris (Fig. 9b). We suspect that such seasonal effects are also the primary reason for the positive rectification signal in northeastern Germany and northern Poland. In southern Europe, these seasonal covariations tend to lead to a negative fossil-fuel rectification effect because the emissions peak in summer (Fig. 2b), when the PBL height is at its seasonal maximum.

The magnitude of the fossil-fuel rectification effect is smaller than the rectifier effect induced by the exchange fluxes with the terrestrial biosphere (Zhang et al., 2016) but still quite substantial. Thus, the fossil-fuel rectification effect clearly needs to be taken into consideration when modeling the atmospheric fossil-fuel CO$_2$ signal, highlighting the need to use and apply accurate time functions. Our results thus clearly support the results of Nassar et al. (2013), who demonstrated the substantial impact of the consideration of time-varying emissions on atmospheric CO$_2$. We extend their result by demonstrating an effect on the annual mean fossil-fuel CO$_2$, suggesting that special attention needs to be given to the relative timing of variations in atmospheric transport and mixing and fossil-fuel emissions. Our results confirm the recent findings by Zhang et al. (2016), who demonstrated the fossil-fuel rectification effect for the first time in a global model. Their signal is locally smaller than ours due to their
Figure 10. Maps of the annual mean fossil-fuel CO₂ signal generated by different countries and regions. (a) Surface pattern created by the emissions from Germany. Panel (b) is as panel (a) but for the France. (c) Column-averaged pattern created by the emissions from Germany. Panel (d) is as panel (c) but for France. Shown are the results for the period 27 March 2008 through 26 March 2009.

use of a much coarser-resolution model, but they also show that the sign of the fossil-fuel rectification effect tends to vary between timescales, with the diurnal being primarily negative, while the seasonal rectification effect being positive. This supports our explanation for the positive signals in northeastern Germany and northern Poland.

5.2 Fossil-fuel CO₂ signal from different sources

Near the surface, the fossil-fuel emissions from a particular region create a distribution that stays mostly within the region of origin (see Fig. 10a and b). The fossil-fuel CO₂ is highly concentrated near the localized areas of high emissions and then drops off quickly by distance with an e-folding spatial scale of a few hundred kilometers. As a result, the fossil-fuel signal tends to be relatively small outside the region of origin, rarely exceeding 1 ppm in contrast to the > 20 ppm signal close to the sources. The different magnitudes of the fossil-fuel CO₂ signals from different regions largely reflects the total emissions, but also the emission intensity, i.e., the emission per unit area. For example, with a total emission of 0.59 Pg CO₂ yr⁻¹, Germany is the biggest source of fossil-fuel CO₂ within Europe, nearly double that of the second-biggest emitter (i.e., France), yet Germany is almost half the size of France, resulting in a considerably higher emission intensity over Germany.

A different picture emerges when considering XCO₂, i.e., the column-averaged dry air mole fraction CO₂. After having been transported aloft, where the fossil-fuel signal can be much more readily dispersed, the imprint of the emissions of any particular region to the fossil-fuel CO₂ within another region is actually quite large (Fig. 10c and d). In a small country, such as Switzerland, only 20% of the fossil-fuel signature in XCO₂ above its territory stems from emissions within, while the contribution of Germany alone is 21% and that of France 18% (Fig. 11). A similar distribution of sources is found for other small countries, such as Austria. In contrast, the fraction of the territorial emissions to the total fossil-fuel signal is quite a bit larger for large countries and regions, such as France or Germany. In the latter case, more than 50% of its total fossil-fuel CO₂ signal stems from emissions within, with four countries contributing most of the remainder. The countries and regions with high overall emissions also contribute, of course, most strongly to the fossil-fuel
CO₂ signal in other countries, with Germany contributing 17 % to the signal in France, 11 % to that in Italy and 28 % to that in the Netherlands. Owing to its lower total emissions, France just contributes 10 % to the signal in Germany and 9 % to that in Italy. Thus, as is the case with classical air pollution, the fossil-fuel CO₂ does not stop at the national borders but extends to continental scales (see Fig. 11).

Among all the processes, the CO₂ emissions from power plants dominate the fossil-fuel distribution, with concentrations reaching up to 16 ppm in the northern part of the domain (see Fig. 12). The point-source nature of this emission sector is clearly visible in the surface distribution, as is the spatially distinct distribution owing to the large differences in power production in the different countries of central Europe. While France has very few fossil-fuel-fired power plants as a result of its high reliance on nuclear and hydroelectric power plants, Germany, Italy, the Netherlands and Poland rely strongly on coal- and gas-fired power plants for their electricity production. This leads to a highly heterogeneous fossil-fuel CO₂ signals of the power plant sector. In total, this sector contributes 32 % to the total fossil-fuel CO₂ signal in central Europe, which is slightly smaller than its contribution to emissions (33 %). This small difference emerges from the somewhat stronger loss of the signal.

Figure 11. Pie charts depicting the origin of the fossil-fuel CO₂ signal for each country/region for the period 27 March 2008 through 26 March 2009. The percentages represent the contribution of each country or region of origin to the total fossil-fuel signal averaged over the air column. The pie chart for Switzerland reveals, for example, that only 20 % of the fossil-fuel CO₂ signal over its territory stems from its territorial emission. Here, CH is Switzerland; DE is Germany; FR is France; IT is Italy; AT is Austria; NL is Netherlands; SW is countries in southwest of the domain; UK is United Kingdom; EA is countries in eastern domain; OT is the rest of countries.
across the lateral boundaries from this sector relative to the signal from the other sectors.

The second-largest fossil-fuel CO$_2$ signal is generated by the emissions from the road transportation sector (22 %) (Fig. 12d), with this share actually being somewhat larger than its share in total emissions (21 %). The transportation sector signal is very smooth due to the distributed nature of the emissions from this sector (see also Fig. 1).

The CO$_2$ signal from the industrial and residential sectors are more granular than that from the transportation sector but still not as distinct as the power plant sector, as there are less country-specific policies impacting the CO$_2$ emissions from these sectors. The emissions and consequently the CO$_2$ signal largely follow population density. The residential sector (mostly heating) contributes 18 % to the total fossil-fuel signal in atmospheric CO$_2$, slightly larger than the emissions from the industrial sector (17 %). These two shares in the signal very nearly reflect their shares in total emissions. The emissions from the “other” sectors (e.g., shipping, waste incineration) is smaller, in comparison (11 %), but not negligible.

The relative contribution of the emissions from the different sectors to the fossil-fuel CO$_2$ vary strongly by region (Fig. 13). Clearly, close to major fossil-fuel-fired power plants, this sector dominates. Owing to the dominance of this mode of electricity production in northern Europe, this signal is particularly strong there. This is most evident over the North Sea, where the advection of the emitted CO$_2$ from the power plants in the UK and the Netherlands creates a particularly visible plume over the ocean. But elsewhere, any of the four major sources can take the leading role. For example, in Switzerland, Paris, and London, the emissions from the residential sector dominate the signal, while over much of southern and western Europe, the transportation sector dominates. The industrial sector dominates the signal in a few hotspot areas, where its emissions are high but no major fossil-fuel-fired power plants are nearby.

These high spatial variations in the relative contribution puts the findings of Vogel et al. (2013) into a spatial context, as they reported for the Heidelberg site a dominance for emissions from power plants (28 %), while the transportation sector contributed only 15 %. This is a typical value for much of western Germany, reflecting the relative contribution of

**Figure 12.** Maps of the annual mean surface fossil-fuel CO$_2$ stemming from different sectors in units of ppm: (a) fossil-fuel-fired power plants, (b) residential heating, (c) industrial processes and (d) road transportation. Shown are the results for the period 27 March 2008 through 26 March 2009.
the different emission sectors (see also Fig. 1). But the contributions are very different, for example, for the CarboCount CH sites in Switzerland (Oney et al., 2015). At Beromünster, the transportation sector dominates over the other sectors, with nearly 70% stemming from this sector alone, while the contribution from power plant emissions is very low at this site, since Switzerland does not operate any fossil-fuel power plants.

These large differences in the relative contribution from the different emission sectors have major implications for the analysis of the fossil-fuel CO$_2$ and how it may change in response to mitigation measures. For example, these large differences will lead to substantial spatial gradients in the CO to CO$_2$ ratio in the fossil-fuel signal, as the different emission sectors have very different CO to CO$_2$ emission ratios. Since CO is often used to identify the fossil-fuel component from atmospheric CO$_2$ observations, these variations need to be carefully disentangled in order to properly diagnose the fossil-fuel component. The strong variations in the contributions from the different sectors thus add a substantial amount of uncertainty to the CO method (Oney et al., 2017; Vardag et al., 2015). A second consequence concerns the detection of changes in emissions from the different sectors. Thus, with the transportation sector contributing little to the very large fossil-fuel signal in much of the northeastern part of our domain, reductions in this sector will be difficult to discern in that region. In contrast, the high relative contribution of the transportation sector to the total signal in southwestern Europe makes it actually quite feasible to detect mitigation measures in this sector in that part of Europe, even though the overall signal might not be that high.

An important caveat of our simulations is the fact that the effective height of the emissions above surface was not considered, but rather all CO$_2$ was released into the lowest model level. As a consequence, the surface CO$_2$ signals from elevated stack emissions from power plants and residential heating are likely biased high relative to those from the transportation sector. Given the large contribution from power plant emissions, it will be important to accurately consider the effective emission height (including plume rise) in future simulations, a point that was also raised by Vogel et al. (2013).
Y. Liu et al.: Fossil-fuel signal

5.3 The response of atmospheric CO$_2$ to an emission reduction

According to their intended nationally determined contributions filed with the United Nations Framework Convention on Climate Change (UNFCCC) in late 2015, the European Union and its member states have agreed to a binding target of a domestic reduction in greenhouse gas emissions of at least 40% by 2030 compared to 1990 (http://www4.unfccc.int/Submissions/INDC/PublishedDocuments/Latvia/1/LV-03-06-EUINDC.pdf). A major question driving international policy making is to what degree such a reduction can be verified through independent means, such as through the monitoring of atmospheric CO$_2$ (Ciais et al., 2014, 2015). To address this question, we conducted several sensitivity experiments to investigate how various reductions in the magnitude and types of emissions affect not only the annual mean fossil-fuel CO$_2$ signal but also its variability. The goal is to determine whether reduced fossil-fuel emissions might be detectable by current and future observing systems, especially satellites. As the satellites have a typical overpass time of 13:00 UTC, we conducted all subsequent analyses using the model data only from this time slot.

Since CO$_2$ is a conservative tracer in the atmosphere on the timescales considered here, a uniform reduction in the emissions leads to a uniform and directly proportional reduction of its current distribution; i.e., a 30% reduction of total fossil-fuel emission would simply lead to a 30% reduction of the fossil-fuel CO$_2$ signal at the surface (Fig. 4a) and throughout the atmospheric column (Fig. 6a). Concretely, the fossil-fuel CO$_2$ would be reduced by more than 4 ppm near the surface for vast stretches of central and northern Europe, with maximum reductions of 10 ppm or more in the emission hotspots (Fig. 14a). This contrasts with the reduction in the column-averaged annual mean XCO$_2$, amounting to just over 0.2 ppm in the regions where the surface decreases by 4 ppm or more (Fig. 14b). A reduction of 0.5 ppm is reached in just a few isolated locations, generally characterized by a high density of point sources, primarily fossil-fuel-fired power plants. Thus, given current measurement accuracies of better than 0.1 ppm for a ground-based observing network (Zellweger et al., 2016), a 30% reduction in the fossil-fuel emissions is fundamentally easily detectable for such a system, although

www.atmos-chem-phys.net/17/14145/2017/ Atmos. Chem. Phys., 17, 14145–14169, 2017
one needs to bear in mind the nontrivial task of separating
the signal from the background variability. In contrast, such a re-
duction in the fossil-fuel emissions is not trivial to detect by
satellite observations for most regions (except around the big
power plants) as it is very challenging to obtain and maintain
accuracies better than 0.5 ppm by current space-based ob-
serving systems (Buchwitz et al., 2015). Furthermore, such
high accuracies are only achieved when the data are aver-
gaged over large scales, i.e., on the order of 1000 km or more.
Nevertheless, taking 0.5 ppm as the threshold for detection
within a single pixel, a 30 % reduction in fossil-fuel emis-
sions thus appears to be beyond the detectability except for a
few hotspot regions (Fig. 14b). Even a 50 % reduction would
not be trivial to detect for a satellite-based system on the ba-
sis of changes in the column-averaged dry air mole fraction.

Given these challenges, a potentially attractive second av-

denue for determining changes in fossil-fuel emissions is the
reduction in temporal variability of atmospheric CO$_2$ that
goes alongside the reduction in the mean signal. This is par-

ticularly promising given the high contribution of the
fossil-fuel CO$_2$ signal to the variability in atmospheric CO$_2$
(see Fig. 8). As is the case for the mean, the conservative
nature of atmospheric CO$_2$ implies that a uniform reduction
of the emissions will lead to a uniform and proportional re-
duction of the variability of the fossil-fuel signal as well.
However, this is not the case for the variability in total at-
mospheric CO$_2$, since covariations between the fossil-fuel
signal and the signal from, for example, the terrestrial bio-
sphere can lead to nonlinear effects. For example, a negative
correlation between the two components would lead to a sit-
uation where the variability of atmospheric CO$_2$ was smaller
than that of the individual components. In such a case, a re-
duction of the fossil-fuel emission would lead to a smaller
decrease in variability than expected. If the two components
were positively correlated, the opposite would occur, i.e.,
the variability in atmospheric CO$_2$ would decrease more than ex-
pected.

Near the surface, the reduction in the temporal standard
deviation and in the mean have nearly the same amplitude for
most places (Fig. 14c). This makes the analysis of changes in
the temporal variability indeed an attractive option to en-
hance the detectability of changes in fossil-fuel emissions.
This is much less the case for the annual mean XCO$_2$, where
the standard deviation changes are in general much smaller
than the changes in the mean, with just a few isolated places
revealing changes in the standard deviation of 0.5 ppm or
more that might be discerned by the current generation of
satellites.

But in these isolated places, the analysis of the temporal
variability might be an interesting option even for satellite-
based measurement systems (Fig. 15). In those places, indi-
cated by the green circles in (Fig. 14c), the changes in the
temporal standard deviation are very large. Even for changes
in emissions of around only 30 %, the changes would be de-
tectable for current satellites (Fig. 15). But the number of
such sites is very low across Europe, making this a special-
ized, rather than general, option.

The detection challenge is not simpler for other potential
emission reduction scenarios, as outlined, for example, in the
EU road map (http://ec.europa.eu/clima/policies/strategies/
2050/index_en.htm). A 50 % reduction in the emissions from
power plants alone (representing a reduction of the overall
emissions by 16 %) results in the mean surface concentration
of atmospheric CO$_2$ going down by more than 2 ppm over
large parts of northwestern Europe, following the pattern of
the surface signal of this sector (see Fig. 12a). In addition,
we find a substantial reduction of the standard deviation of surface atmospheric CO\textsubscript{2} by more than 2 ppm in these regions, with the hotspots of power plant emissions showing a reduction in the standard deviation of atmospheric CO\textsubscript{2} of 5 ppm or more. The reduction of the average annual mean column XCO\textsubscript{2} is much smaller than that of atmospheric CO\textsubscript{2} at the surface, amounting to little more than 0.2 ppm over wide swaths of northern Europe. The maximum reductions are of the order of 0.5 ppm in the proximity of large clusters of fossil-fuel-fired power plants, i.e., generally too small to detect. But, in these regions, the changes in the variability in XCO\textsubscript{2} is quite high, making this method again potentially attractive for detecting changes. In fact, in several regions, including some major cities, a 19 % reduction of the fossil-fuel emissions would result in a change of more than 0.5 ppm in the standard deviation, i.e., above detection level. This thus supports the findings of Pillai et al. (2016) that changes in fossil-fuel emissions are fundamentally detectable over major cities or major point sources, but it also shows that this detection is very challenging.

The signals get even more difficult to discern if the emission reductions occur in individual sectors other than the power plants. For example, detectable signals by current generation satellites occur only if industrial emissions are cut by more than 80 % or if residential emissions are cut by more than 90 %. Also country-level emissions are not trivial to be clearly detected. A reduction in Germany by 50 % is potentially detectable by current satellites, with a maximum reduction of XCO\textsubscript{2} by 0.95 ppm. For most other countries, however, a 50 % reduction in emissions is difficult to detect.

All the analyses here relied on using the model output on all available days; i.e., we assumed perfect temporal coverage. This is overly optimistic, since cloud cover and other complicating factors (e.g., aerosol layers) will cause the coverage to decrease considerably, complicating the detection. We assumed here also “perfect transport”, i.e., no errors in how the emission reductions manifest themselves in a change in the concentration field. In fact, errors in this transport are perhaps, besides the lack of observations, the largest impediment to detect changes in fossil-fuel emissions.

But regardless of this additional challenge, there is much additional information contained in high-frequency observations of atmospheric CO\textsubscript{2}. As we demonstrated above, the temporal variations are potentially highly useful for detecting fossil-fuel emissions changes from various sources, especially those with a strong spatial granularity such as power plants or individual cities. For a routine monitoring of strong point sources, Velazco et al. (2011) therefore proposed a constellation of five satellites that combine imaging capability with a relatively wide swath (Bovensmann et al., 2010). Such a constellation would offer daily global coverage, though the presence of clouds would reduce the effective coverage considerably. As the precision and accuracy of satellite retrieved XCO\textsubscript{2} will improve in the future, that minimum change will go down as well.

6 Summary and conclusions

We have investigated the fossil-fuel signal in atmospheric CO\textsubscript{2} over southern and central Europe using a regional high-resolution atmospheric model forced with temporally and spatially highly resolved variations in the fossil-fuel emissions. The assessment of the modeled atmospheric CO\textsubscript{2} with in situ measurements on the highest level across multiple sites across Europe reveals good agreement on all timescales considered with biases of less than 1.5 ppm, with the exception of the tall tower site Hegyhatal in central Hungary. The model is also able to capture the reconstructed fossil-fuel component at two sites quite successfully. Although the model tends to underestimate the amplitude of the daily-averaged fossil-fuel CO\textsubscript{2} in winter, the simulation matches fossil-fuel CO\textsubscript{2} from both sites very well most of the time, revealing the high quality of the transport model and reasonable time profiles of the fossil-fuel emissions used as input.

Over much of Europe, the fossil-fuel CO\textsubscript{2} is a dominant component of the spatial variability of atmospheric CO\textsubscript{2}, particularly near the surface. In some places, it even contributes significantly to the total (including background) CO\textsubscript{2}, particularly in large urban centers and along power plant plumes. Also the contribution to the temporal variability is very substantial. Fossil-fuel CO\textsubscript{2} makes a particularly large contribution on synoptic and diurnal timescales whereas the seasonal variability is dominated by biospheric activity. The influence is large not only over the hotspot regions of fossil-fuel emissions but also over large areas downstream. In case of diurnal variability, fossil-fuel CO\textsubscript{2} is the dominant component over wide areas of northern and western Europe.

Temporal variability of the emissions has a non-negligible influence on annual mean fossil-fuel CO\textsubscript{2} mole fractions near the surface due to diurnal and seasonal rectifier effects. Differences between annual mean values with temporally variable and constant emissions can be up to a few ppm in the hotspot regions but are mostly below 1 ppm elsewhere. This implies that temporal variability of fossil-fuel emissions needs to be accurately represented for realistic simulations, confirming the results of Zhang et al. (2016). It is also important for reliably detecting fossil-fuel emission changes from specific sources since different sources have different temporal profiles.

Simulating fossil-fuel emissions from different countries and sectors suggests that the major part of the signal near the surface remains in the country of origin. Ground-based in situ observations are thus most sensitive to fossil-fuel emissions from the country where they are located. A different picture emerges for column-averaged dry air mole fractions (XCO\textsubscript{2}) as measured by satellites, for which the signal is much more dispersed. Only over Germany is the contribution from emissions within the country larger than 50 %, whereas over France the signal from neighboring countries dominates (66 %). An important reason for these contrasting results seems to be the differences in electricity produc-
tion, which mostly relies on nuclear power in France but on fossil fuels in its neighboring countries including Germany, UK and Italy. Over small countries such as Switzerland or the Netherlands, the contribution from abroad is typically the dominating component. Among all the processes, fossil-fuel emissions from power plants contributes the most (approx. one-third) to the total fossil-fuel signal of CO\(_2\) both at the surface and in the column. However, the power plant signal at the surface is likely overestimated in our simulations, since all emissions were released into the lowest model level without considering the true elevation of the source. The signal from power plant emissions has a pronounced and distinct spatial pattern that provides us an opportunity to discern changes in from power plant emissions from changes in other sources.

Based on a number of sensitivity studies, we show that reductions in fossil-fuel emissions leave a distinct signal not only in the time mean distribution of atmospheric CO\(_2\) but also in its temporal variability. This opens potentially additional ways to detect and verify emission reductions. But this opportunity exists primarily for surface-based measurement networks, while the satellite-based systems that measure the column-averaged XCO\(_2\) will see too small changes, in general, relative to their current measurement capabilities. An important exception are a few hotspot sites, where the satellites will be able to detect fairly modest changes of about 30 % when assuming an accuracy of the satellite observations of 0.5 ppm.

As both satellite and surface measurements have advantages and disadvantages, combining surface measurements with satellite data and increasing the frequency and coverage of the latter will be the optimal path forward to enhance the possibility of detecting future changes in fossil-fuel emissions.

Data availability. The model output underlying this publication is available through the ETH library (https://www.research-collection.ethz.ch) under the https://doi.org/10.3929/ethz-b-000213651.

The Supplement related to this article is available online at https://doi.org/10.5194/acp-17-14145-2017-supplement.

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

Acknowledgements. This study was funded by the Swiss National Foundation (SNF) as part of the CarboCount CH Sinergia Project (grant number CRSII2 136273). Additional funding was provided by ETH Zürich (Yu Liu and Nicolas Gruber) and Empa (Dominik Brunner). We thank the Center for Climate Systems Modeling (C2SM) and especially Anne Roches for providing extensive support. We acknowledge MeteoSwiss for the provision of their operational COSMO analysis products. The computations were done on the supercomputers of the Swiss National Supercomputing Centre (CSCS). We are very thankful to the principal investigators responsible for the Mace Head (MHD) station in Ireland, for the SNO-ICOS-France station Puy de Dôme (PUY) in France, for the Hegyhatsal station (HUN) in Hungary, and for the Cabauw station (CBW) in the Netherlands for making their data publically available. We are also grateful to Sander van Der Laan and Felix Vogel for sharing their estimates of the fossil-fuel component at Lutjewad and Heidelberg, respectively. Furthermore, we are very grateful to Woutler Peters, Wageningen University, for providing the CO\(_2\) global model boundary conditions and to Christoph Gerbig for providing the VPRM biospheric flux data. Finally, we acknowledge the comments and suggestions of the two reviewers, whose input helped us to improve the paper.

Edited by: Rolf Müller
Reviewed by: two anonymous referees

References


Kuenen, J. J. P., Visschedijk, A. J. H., Joziwcka, M., and Denier Van Der Gon, H. A. C.: TNO-MACC-II emission inven-


Pillai, D., Buchwitz, M., Gerbig, C., Koch, T., Reuter, M., Bovensmann, H., Marshall, J., and Burrows, J. P.: Tracking city CO2 emissions from space using a high-resolution inverse mod-


