Sensitivity of free tropospheric carbon monoxide to atmospheric weather states and their persistency: an observational assessment over the Nordic countries

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Abstract. Among various factors that influence the long-range transport of pollutants in the free troposphere (FT), the prevailing atmospheric weather states probably play the most important role in governing characteristics and efficacy of such transport. The weather states, such as a particular wind pattern, cyclonic or anticyclonic conditions, and their degree of persistency determine the spatio-temporal distribution and the final fate of the pollutants. This is especially true in the case of Nordic countries, where baroclinic disturbances and associated weather fronts primarily regulate local meteorology, in contrast to the lower latitudes where a convective paradigm plays a similarly important role. Furthermore, the long-range transport of pollutants in the FT has significant contribution to the total column burden over the Nordic countries. However, there is insufficient knowledge on the large-scale co-variability of pollutants in the FT and atmospheric weather states based solely on observational data over this region. The present study attempts to quantify and understand this statistical co-variability while providing relevant meteorological background.

To that end, we select eight weather states that predominantly occur over the Nordic countries and three periods of their persistency (3 days, 5 days, and 7 days), thus providing in total 24 cases to investigate sensitivity of free tropospheric carbon monoxide, an ideal tracer for studying pollutant transport, to these selected weather states. The eight states include four dominant wind directions (namely, NW, NE, SE and SW), cyclonic and anticyclonic conditions, and the enhanced positive and negative phases of the North Atlantic Oscillation (NAO). For our sensitivity analysis, we use recently released Version 6 retrievals of CO at 500 hPa from the Atmospheric Infrared Sounder (AIRS) onboard Aqua satellite covering the 11-year period from September 2002 through August 2013 and winds from the ECMWF’s ERA-Interim project to classify weather states for the same 11-year period.

We show that, among the various weather states studied here, southeasterly winds lead to highest observed CO anomalies (up to +8 %) over the Nordic countries while transporting pollution from the central and eastern parts of Europe. The second (up to +4 %) and third highest (up to +2.5 %) CO anomalies are observed when winds are northwesterly (facilitating inter-continental transport from polluted North American regions) and during the enhanced positive phase of the NAO respectively. Higher than normal CO anomalies are observed during anticyclonic conditions (up to +1 %) compared to cyclonic conditions. The cleanest conditions are observed when winds are northeasterly and during the enhanced negative phases of the NAO, when relatively clean Arctic air masses are transported over the Nordic regions in the both cases. In the case of nearly all weather states, the CO anomalies consistently continue to increase or decrease as the degree of persistency of a weather state is increased. The results of this sensitivity study further provide an observational basis for the process-oriented evaluation of chemistry transport models, especially with regard to the representation of large-scale coupling of chemistry and local weather states and its role in the long-range transport of pollutants in such models.

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1 Introduction

Apart from the local sources of pollution that degrade local air quality and hence human health, many studies show that, depending on the global and regional circulation patterns and favourable meteorological conditions, the long-range transport of pollutants also contributes to increased pollutant concentrations. In fact, the importance of hemispheric and long-range transport of pollutants is now widely recognized in the scientific community, and the research focus in recent years has deservedly been on better characterization of source-to-sink relationships and drivers of pollutant variability during such transport (Li et al., 2002, 2005; Stohl et al., 2002; Creilson et al., 2003; Eckhardt et al., 2003; Trickl et al., 2003; Duncan and Bay, 2004; Pfister et al., 2004; Huntrieser et al., 2005; Chin et al., 2007; Shindell et al., 2008; Fiore et al., 2009; Dentener et al., 2010; Brandt et al., 2012; Christoudias et al., 2012; Lin et al., 2012). It is important to keep in mind that it is the local meteorology and synoptic-scale weather patterns that eventually determine the spatio-temporal distribution of pollutants, their transport characteristics and final fate. One of the main mechanisms by which the pollutants (e.g. from wildfire emissions) get transported from their source regions to the Earth’s northernmost latitudes as far as the Arctic is through varying states of atmospheric circulation. This was first realized through the phenomenon of Arctic haze observed during the winter/spring months that was first reported in the 1950s (Quinn et al., 2007). It is now known that the major pathways to the Arctic depend upon the season and the position of the Arctic front. For example, during the winter and spring months, the intense Siberian high-pressure system pushes the Arctic front towards the south whereby the polluted regions of the Eurasian subcontinent are within the Arctic air mass resulting in the efficient transport of pollutants during this time of the year.

The spatio-temporal distribution of pollutants over the Nordic countries is a result of complex interplay of local sources, atmospheric circulation patterns, and contributions from the long-range transport originating from North America, continental Europe and Asia. The dominant modes of atmospheric variability in the Northern Hemisphere affecting the Nordic countries, especially in winter, are the North Atlantic Oscillation (NAO) and Arctic Oscillation (AO). The positive and negative phases of NAO are marked by changes in the wind speed and direction over the Atlantic, heat and moisture transport across the Atlantic. The frequency and intensity of the number of storms and warm conveyor belts influence the transatlantic transport of pollutants from North America to Europe, including over the northern European latitudes (Li et al., 2002; Hurrel et al., 2003; Duncan and Bey, 2004; Dentener et al., 2010). Eckhardt et al. (2003) observed a strong correlation between the NAO and transport of anthropogenic pollution into the Arctic and eastern Europe from all the Northern Hemisphere continents with an enhanced transport during the positive NAO phase. However, a significant anti-correlation is observed between NAO and the anthropogenic pollutants over western and central Europe (Christoudias et al., 2012). Significant correlations between the positive phase of AO and elevated ozone concentrations in western Europe is observed and can be attributed largely to in situ production associated with the subsidence within the high-pressure dome or entrainment of pollutants into this dome (Creilson et al., 2003). Pfister et al. (2004) showed that when averaged over Europe, almost 67% of the anthropogenic carbon monoxide (CO) at the surface comes from regional sources, in addition to the transport from North America (14%) and Asia (15%). However, at higher altitudes, the contribution from North America and Asia is significantly higher. Brandt et al. (2012) using the 3-D long-range chemistry transport hemispheric model showed that the contributions from North American anthropogenic emissions to the ozone levels in European subcontinent is 3.1% and the contributions from European anthropogenic sources to North America is 0.9%.

One of the major pathways carrying pollutants from the continental Europe to the Arctic passes over the Nordic countries. Tang et al. (2009) studied the long-range transport and weather patterns relating to high ozone events in southern Sweden, and using a trajectory model showed that these events occurred during anticyclonic events, especially during summer. But they observed strong negative relationship between cyclonic and high ozone events. Recently, Devasthale and Thomas (2012) investigated co-variation of temperature inversions and CO over Scandinavia during winter using satellite sensor data. They showed that the increased levels of CO are observed when the atmosphere is thermodynamically unstable (weaker or negative inversion strength) and when the westerly winds are strong. Apart from long-range transport of pollutants in the free troposphere (FT), in cold climate of the Nordic countries, unfavourable meteorological conditions such as thermal inversions, low boundary layer height and low temperatures can contribute to increased pollutant concentrations in the lowermost troposphere near the surface. The above-mentioned studies emphasize the need for a better quantification of the linkages between the pollutant concentrations and atmospheric weather states.

Many of the studies mentioned above examine CO, since CO is often considered as an excellent tracer to investigate pollution transport characteristics due to its moderate lifetime in the atmosphere. Increased carbon monoxide levels would not only enhance carbon dioxide levels in the atmosphere through its reaction with hydroxyl (OH) radicals, but also indirectly increase concentrations of short-lived climate pollutants such as ozone and methane, which would otherwise be depleted by OH radicals. Therefore, monitoring CO and understanding its sensitivity to large-scale weather patterns, based solely on observations, is important not only to gain insights into long-range pollution transport, but also to serve as an observational basis for the sensitivity studies to evaluate chemistry transport models.
In spite of their importance as mentioned above, there is no consistent observationally based assessment of how the dominant weather states impact FT CO variability over the Nordic countries. The present study attempts to partially fill this gap. Decreased instrument sensitivity over very cold surfaces, variable snow cover and difficulties in cloud detection are some of the factors that limit the use of satellite remote sensing to study the atmospheric composition variability over the Nordic regions, especially in the lowermost troposphere. But the data records of atmospheric composition from satellite sensors, especially from hyperspectral sounders such as IASI (Clerbaux et al., 2009) and AIRS (Chahine et al., 2006), are continuously improving and we now have better understanding of their retrieval quality and sensitivities. More than a decade long data, e.g. from AIRS and MOPITT, can be exploited to investigate statistics on the large-scale co-variability of weather states and trace gases, as attempted here.

In the next section, we describe the data set used and methodology adopted, followed by presentation of an overview of dominant atmospheric circulation patterns and corresponding meteorological conditions over the Nordic region, with specific focus on Sweden, in Sect. 3. We then discuss sensitivity of CO to these patterns and persistency of these patterns in Sect. 4. The final section presents conclusions.

2 The Atmospheric Infrared Sounder (AIRS) CO and ERA-Interim data sets

AIRS onboard Aqua satellite has 2378 hyperspectral channels, out of which about 36 well-defined channels with wavenumbers ranging from 2181.49 to 2221.12 cm\(^{-1}\) are used in V6 to retrieve CO (http://disc.sci.gsfc.nasa.gov/AIRS/documentation/v6_docs/v6releasedocs-1/V6_Retrieval_Channel_Sets.pdf). A priori profiles (sets of 100 layers) with monthly granularity are used as a first guess. The profiles are based on MOZART (Model for OZone And Related chemical Tracers) monthly mean hemispheric profiles (http://disc.sci.gsfc.nasa.gov/AIRS/documentation/v6_docs/v6releasedocs-1/V6_CO_Initial_Guess_Profiles.pdf). By varying the geophysical state, the retrieval algorithm for CO basically tries to minimize the weighted difference between clear-sky radiance and the radiance computed using a forward model. Using averaging kernels, the retrieval algorithm relates estimated CO profile to “true” profile and a priori information. The algorithm details are described in Susskind et al. (2003) and in Warner et al. (2007, 2010, and 2013).

We use the recently released Version 6, daily standard and support Level 3 retrievals of CO (AIRS Science Team/Joao Texeira, 2013; AIRS-V6L3UG, 2013). Eleven years of data from September 2002 to (and including) August 2013 are analysed. The AIRS retrievals of temperature and CO are validated and matured considerably over the years to enable variability studies (Divakarla et al., 2006; Fetzer, 2006; Warner et al., 2007, 2010, 2013; Yurganov et al., 2008). The accuracy and biases of AIRS CO are well documented in the studies mentioned above.

Since the focus of the present study is on the FT, we have analysed CO at four different vertical levels, namely 850 hPa, 700 hPa, 500 hPa and 400 hPa, but the results are shown only for 500 hPa. The reasons for that are (a) the signal of pollutant transport in the FT is most tangible at this level, (b) coincidentally, AIRS retrievals are of best quality at this level (Yurganov et al., 2008; Warner et al., 2010), and (c) for the sake of brevity. The tendencies in CO observed at these four vertical levels and corresponding wind patterns during selected weather states are not significantly different, since many of the weather states, when they are persistent, affect the entire FT (as shown later).

We analyse AIRS retrievals only when the degrees of freedom value is larger than 0.5. For investigating large-scale features and tendencies, as attempted in the present study, AIRS Level 3 CO data are quite suitable, as for example shown by Devasthale and Thomas (2012). Data from both ascending and descending passes of the afternoon Aqua satellite are used. We have allowed up to 30 % cloud cover while analysing the AIRS CO retrievals based on the findings of our sensitivity studies (Devasthale and Thomas, 2012) and previous experience with AIRS data (Devasthale et al., 2010, 2011, 2012, 2013). Susskind et al. (2003) have previously presented detailed analysis of the accuracy of AIRS retrievals in the presence of clouds. The yield and accuracy of AIRS retrievals should not degrade significantly up to 30 % cloud cover. Recently, Warner et al. (2013) showed that the AIRS CO retrievals in cloud-contaminated cases are of comparable quality. The degrees of freedom of the signal, an indicator of information content, are reduced only by up to 0.2 in cloudy cases (please refer Figs. 3 and 4 in Warner et al., 2013). The difference should even be smaller in our cases, since we allow only 30 % cloud contamination. Furthermore, the majority of opaque clouds occurring over the study area are low clouds (cloud tops less than 700 hPa). Since we analysed retrievals at 500 hPa, the cloud impact is estimated to be small. Finally, the absence of any spatial correlation between cloud fraction and observed CO anomalies also suggests that the cloud impact is negligible.

The advantage of using AIRS data lies in the fact that (a) the simultaneous retrievals of temperature and humidity in time and space are available which can be used to understand thermodynamical properties of the atmosphere and possible transport of heat and moisture during different weather states, (b) the longest (> 11 year) data record of CO from hyperspectral measurements is available, and (c) the synergy with other A-Train sensors providing aerosol and cloud information can be exploited in future studies.
For investigating winds, we used 6-hourly zonal \( (u) \) and meridional \( (v) \) wind components from the ECMWF’s ERA-Interim reanalysis (Dee et al., 2011) for the same period when AIRS CO data are available.

### 3 An overview of selected weather states

In the present study, we select eight weather states that most frequently occur over the study area (42–80° N, 10° W–40° E). Figures 1–5 show an overview of circulation patterns and typical meteorological conditions observed under these weather states. The states are selected based on the synthesis of previous literature (e.g. Chen, 2000; Linderson, 2001) and further confirmed by manual inspection of numerous weather reports from the Swedish Meteorological and Hydrological Institute. Since the persistency of a weather state may enhance or reduce pollution levels in the FT, for each selected weather state, we have also investigated tendencies of CO anomalies under three persistency periods, namely 3 days (P3), 5 days (P5) and 7 days (P7). For brevity, we present the circulation patterns and meteorological conditions only for the P5 case, but the sensitivity results for CO are shown for all weather states and persistency periods later in this study.

The eight identified weather states consist of four dominant wind directions (NW, NE, SE and SW), anticyclonic and cyclonic conditions, and two enhanced phases of the NAO. In the case of the first four weather states, we chose the centre (55–60° N, 12–20° E) of the study area (45–80° N, 10° W–40° E) to average daily wind speed and direction at 850 hPa from the ERA-Interim reanalysis. Based on these daily averages we selected days when a particular wind direction prevailed and persisted for at least 3, 5 and 7 days. The same procedure is applied for selecting anticyclonic and cyclonic conditions based on average mean sea level pressure (MSLP) over the centre of the study region. For the remaining two weather states, the selection of days is based on NAO indices. The overlapping dates among weather states are intuitively avoided by the algorithm. For example, if a certain day is assigned to a certain weather state, then that day is not considered further in the statistics of other weather states. But the selected days are inclusive within the persistency periods of the same weather state. For example, when a weather state persists for 7 consecutive days, then the first three and five days of such event are included in the corresponding P3 and P5 cases. We analysed weather state data in the following order. First, the time information for P3, P5, and P7 for the NAO cases are obtained, followed by the analysis for anticyclonic and cyclonic conditions and finally for the wind directions (clockwise NW, NE, SE, SW).

During the selected 11-year study period, relatively speaking, these states (i.e. NW, NE, SE, SW, anticyclonic and cyclonic, and EP and EN) occur 9, 3, 4, 14, 28, 27, 6 and 9% of the time respectively. The number of events studied for each state is mentioned in Table 1. The probability of a particular weather state prevailing over the study area decreases with increasing persistency. Consequently, the results for 7-day periods, shown later, are in some cases patchy. However, the CO anomalies exceed at least one standard deviation and hence are significant.

<table>
<thead>
<tr>
<th>P3</th>
<th>P5</th>
<th>P7</th>
</tr>
</thead>
<tbody>
<tr>
<td>NW</td>
<td>180</td>
<td>72</td>
</tr>
<tr>
<td>NE</td>
<td>85</td>
<td>31</td>
</tr>
<tr>
<td>SE</td>
<td>63</td>
<td>25</td>
</tr>
<tr>
<td>SW</td>
<td>280</td>
<td>100</td>
</tr>
<tr>
<td>Anticyclonic</td>
<td>556</td>
<td>224</td>
</tr>
<tr>
<td>Cyclonic</td>
<td>540</td>
<td>218</td>
</tr>
<tr>
<td>EP</td>
<td>121</td>
<td>48</td>
</tr>
<tr>
<td>EN</td>
<td>178</td>
<td>72</td>
</tr>
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</table>

Another important characteristic of the Earth’s atmosphere is pressure distribution as it defines the wind and weather patterns globally. Figure 3 shows the composites of the magnitude and wind direction at 850 hPa during high MSLP conditions and low MSLP conditions over the centre of the study. During high MSLP conditions, the winds seem to favour the transatlantic transport towards the northernmost latitudes.
Figure 1. Atmospheric circulation patterns at 850 hPa when winds (in m s\(^{-1}\)) are NW and NE over the centre of the study area. The colourbar indicates wind strength (in m s\(^{-1}\)). The study area is marked with black rectangle.

The anticyclonic flow further circulates air masses from continental Europe to over the Norwegian Sea and the northern parts of the study area. On the other hand, during low MSLP conditions, the winds have a much lower trajectory in the Atlantic and the air masses advected from across the Atlantic are transported over continental Europe and are caught up in the cyclonic flow centred around central Scandinavia. The circulation pattern during anticyclonic (cyclonic) conditions leads to enhanced (reduced) heat and moisture transport over the western part of Scandinavia and Northeast Atlantic Ocean as shown in Fig. 5.

The gradients in pressure and hence, the winds, force different types of oscillations. One such prominent oscillation, manifested in boreal winter as a see-saw in pressure over the Atlantic, is the North Atlantic Oscillation. As described in the Introduction, the NAO phases play an important role in the transatlantic transport of pollutants. Shown in Fig. 4 are respectively the 850 hPa winds associated with enhanced positive (EP: NAO index > +1) and enhanced negative NAO conditions (EN: NAO index < −1) and when these conditions prevail for at least five consecutive days. The daily NAO index for the period in study was downloaded from the following link: http://www.cpc.ncep.noaa.gov/products/precip/CWlink/pna/nao.shtml. The NAO index itself does not show any significant trend during the last decade as shown in the Supplement Fig. S1.

During the enhanced positive NAO phase, the winds are stronger and there is a significant advection of air masses across the Atlantic from northern US and Canada into northeastern Europe and Scandinavia (Fig. 4a). In fact, there is striking resemblance between warmer temperature anomalies (Fig. 5) and wind pattern in the EP case, suggesting the efficiency of atmospheric transport. During the EN phase, the winds are much weaker (Fig. 4b) and the cold Arctic air masses propagate into Scandinavian countries (also clearly visible in Fig. 5) and there is a relatively stronger southwesterly flow over northern Europe.

The normalized frequency of number of days of data available for each weather state as function of months is shown.
in Fig. 6. The distribution of their occurrence is not always uniform as expected, since different weather states are dominant during different times of a year except for anticyclonic and cyclonic periods (i.e. during above normal and below normal MSLP conditions) when their occurrences are distributed evenly. The frequency distribution of EP and EN phases is such that the enhanced positive NAO phases are more prominent during the winter half of the year, while the enhanced negative NAO phases are more prominent during the other half of the year. This unequal distribution of samples as a function of months makes it difficult to compare their relative impact of weather states on observed CO levels due to interference of seasonality of CO. To address this, we calculate 11-year annual climatology of CO by taking a weighted average based on the distribution of occurrence for a particular state as a function of month as follows:

\[ C = \sum_{i=1}^{12} w_i c_i \]  

where \( i \) is month, \( w_i \) is monthly weight (based on figure shown above), \( c_i \) is monthly climatology of CO.

We then subtract this climatology from the composite of CO observed during that state to compute anomalies. This ensures that we remove the seasonal variations while comparing different states and that the observed CO anomalies are indeed due to contribution from that particular weather state and its persistency.

4 Sensitivity of CO to weather states and their persistency

The sources of CO over the study are mainly anthropogenic resulting from fossil fuel burning, vehicular emissions and industrial activities. The biomass burning (natural and anthropogenic) also contributes to the total CO budget. The seasonality in photochemical production and depletion gains importance with increasing altitude. This in combination with long-range and inter-continental transport drives the seasonal variability of CO in the free troposphere over the study area. The climatological seasonal distribution of CO at 500 hPa over the study area is shown in Fig. S2. As expected the CO concentrations are higher in the late winter to early spring due to their increased lifetime as the photochemical loss is at its minimum because of the lack of sunlight, and due to increased emissions and efficient transport during this time of the year.

Figure 7 shows CO anomalies in the free troposphere (500 hPa) for four chosen wind directions and persistency periods of P3, P5 and P7. Only statistically significant anomalies exceeding one standard deviation are shown. When the winds are NW, above normal CO concentrations are observed over northern Europe and CO anomalies increase.
Figure 6. Normalized distribution of the number of weather events as a function of month when they persisted for 5 days for (a) wind directions, (b) anticyclonic and cyclonic cases and (c) for enhanced positive and negative NAO.

Figure 7. Normalized distribution of the number of weather events as a function of month when they persisted for 5 days for wind directions, anticyclonic and cyclonic cases and for enhanced positive and negative NAO.

The deviation of the CO concentrations under anticyclonic and cyclonic conditions and their persistency is shown in Fig. 8. The CO concentrations are in general higher over northern Europe during anticyclonic conditions compared to cyclonic. The CO concentrations continue to increase as anticyclonic conditions persist, and vice versa for cyclonic situations. A careful analysis of wind patterns reveal that, during anticyclonic conditions, the polluted air masses from continental Europe and North America are being drawn and circulated over the Nordic regions whereas, during cyclonic conditions, cleaner Arctic air is mixed in the circulation gyre thereby being more efficient in the removal and dispersal of pollutants resulting in relatively cleaner conditions.

Lastly, the sensitivity of CO to NAO phases and their persistency is shown in Fig. 9. The CO anomalies are higher during the EP phase compared to the EN phase during all persistency periods. Furthermore, there is a clear tendency that, as the positive phases of the NAO persist, CO concentrations tend to increase, especially in the higher latitudes. The free troposphere on the other hand becomes cleaner when the negative phases of NAO persist. When the westerlies are weakened, cold and clean Arctic air is drawn over northern Europe during the negative phase. The tendencies in CO observed during positive and negative phases of NAO are consistent with previous studies that use models simulations (Eckhardt et al., 2003; Christoudias et al., 2012). For example, from the analysis of 15-year simulations, Eckhardt et al. (2003) show enhanced tracer transport to the Arctic that passes over the Nordic countries during positive phases of the NAO. Christoudias et al. (2012) arrive at a similar conclusion with regard to transport towards northern Europe.

To quantify the importance of the different synoptic states, Fig. 10 shows the percentage change in CO in the FT observed during the different weather states and persistency periods over the study area. It can be seen that in nearly all the cases, the CO concentrations either steadily increased or decreased with increased persistency of each weather state. The highest CO contribution, almost 4–8% depending on the degree of persistency, is observed when the winds are SW. The second (up to 4%) and third highest (2.5%) anomalies are observed during NW winds and the enhanced positive phases of the NAO respectively. The CO anomalies of completely opposite signs during positive and negative phases of the NAO confirm the significance of the role of natural variability in pollutant transport and diffusion. The anticyclonic and cyclonic conditions also show opposite signs of CO anomalies, with maximum anomalies in the order of 1% observed during NW winds and the enhanced positive phases of the NAO respectively. The CO anomalies of completely opposite signs during positive and negative phases of the NAO confirm the significance of the role of natural variability in pollutant transport and diffusion.
transport can lead to much higher changes in CO over the study area.

As mentioned in Sect. 2, although we show results of CO variability at 500 hPa, we have investigated this variability at four different levels in the FT and in the total column CO as well. As the persistency period of the chosen weather states increases, it is expected to affect the CO variability in the entire troposphere in a systematic manner. This is evident in Fig. S3 which shows an example of the impact of wind directions on the total column CO variability. The tendencies in CO total column anomalies under different wind directions and across persistency periods are strikingly similar to those observed at 500 hPa. This underscores the importance of the chosen weather states in regulating CO variability in the entire troposphere. Under certain conditions, for example very cold winters and surfaces, the sensitivity and information content of AIRS may peak only in the middle troposphere and the total column values are affected by this problem. But
pollutants from the source regions, it is the local weather of the year due to the presence of inversions over the study region (Devasthale and Thomas, 2012). Then the likelihood of strong episodic vertical injections of pollutants exists only during summer months (via dry or moist convection), but it is very small since such events are usually few in number. And finally, in the FT (850–400 hPa), the atmospheric variability over the study region is governed by the large-scale frontal systems created by the baroclinic disturbances (esp. in the winter half of the year). These systems provide a conducive environment for the advective rather than the local convective transport. Our study area is usually at the receiving end of such systems that often arrive from the neighbouring oceanic areas (Northeast Atlantic, North Sea, or the Arctic Ocean) or from the continental Europe.

The other transport mechanism, i.e. warm conveyor belts, will intrinsically be included in some of the weather states studied here, especially in those cases where winds have history over the North Atlantic. But it should be noted that, irrespective of the mechanism that triggered the transport of pollutants from the source regions, it is the local weather state over our study area that will regulate the distribution of these pollutants. For example, a particular weather state may either dampen the isentropic transport of pollutants or facilitate it further to other regions (e.g. to the Arctic). For investigating the possible impact of warm conveyor belts on the observed pollutant variability, we need to take into account full transport history from the source to the study area. This is currently being addressed in a separate study.

5 Conclusions

Although the long-range transport governs the variability of pollutants in the FT over the Nordic countries, it is the atmospheric weather states that finally determine the spatio-temporal distribution and the fate of the pollutants. The persistency of a particular weather state may further enhance or reduce concentrations of pollutants. Understanding the statistical link between weather states and pollution variability is not only crucial to understand the role of long-range transport itself, but also to be able to simulate such a link in chemistry transport models. The latter is important since chemistry transport models are often used to estimate changes in pollution load, attribution studies and developing mitigation strategies under different climate change scenarios. In this context, the present study attempts to provide insights into the sensitivity of free tropospheric carbon monoxide to different weather states and their degree of persistency based solely on observational data.

We investigated free tropospheric CO variability during eight weather states often prevailing over the Nordic countries. Selected states include four wind directions (NW, NE, SE, and SW), anticyclonic and cyclonic conditions, and positive and negative phases of the NAO. Furthermore, we investigated tendencies in CO under three different degrees of persistency (3-day, 5-day and 7-day) of each weather state. For nearly all the weather states, CO levels consistently continued to increase or decrease as the degree of their persistency increased. Among the weather states studied here, relatively speaking, the highest CO anomalies were observed when winds had southeasterly component, transporting pollutants from the central and eastern European regions to over the Nordic countries. The second largest contribution was from the northwesterly winds, most likely carrying pollutants as a result of long-range transport from polluted North American regions. The third largest anomalies are observed during enhanced positive phase of the North Atlantic Oscillation, confirming the importance of this natural variability in controlling pollutant distribution and transport over the study region. The cleanest conditions were observed under prevailing northeasterly winds and the enhanced negative phase of the NAO. The results from this sensitivity study provide an observational foundation for the process-oriented evaluation of chemistry transport models.
It must be mentioned that although we provide relevant information on atmospheric circulation and meteorology while inferring the potential role of long-range pollution transport in the observed sensitivity of CO to weather states, the actual attribution and precise quantification of contribution from different transport pathways must be done using trajectory or chemistry transport models.

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