Influence of the North Atlantic Oscillation on air pollution transport

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Abstract

We examined the influence of the North Atlantic Oscillation (NAO) on the atmospheric dispersion of pollution by computing the emission, transport and removal of insoluble gaseous and water-soluble aerosol tracers, tagged by the continent of origin. We simulated a period of 50 yr (1960–2010), using the ECHAM/MESSy atmospheric chemistry (EMAC) general circulation model. The model accounts for anthropogenic, biogenic and biomass burning sources, removal of trace gases through OH oxidation, and precipitation, sedimentation and deposition of aerosols. The model is shown to reproduce the observed spatial features of the NAO, moisture transports and precipitation. During high NAO phase seasons the axis of maximum westerly North American trace gas transports extends relatively far to the north and east over Europe. The NAO phase is significantly correlated with North American tracer concentrations over the northwestern Atlantic Ocean and across northern Europe, and with European trace gases and aerosols beyond the arctic circle. Our results indicate marked differences and partly reversed correlations for the insoluble gas and the soluble aerosol tracers. We find a strong anti-correlation over western and central Europe between European pollutant gas and aerosol concentrations and the phase of the NAO.

1 Introduction

The North Atlantic Oscillation (NAO) is a major recurring large-scale pattern of interannual variability in the atmospheric circulation of the Northern Hemisphere. The NAO is manifested by swings in the meridional atmospheric pressure difference between the sub-polar and sub-tropical Atlantic Ocean, most prominently during boreal winter, and is associated with changes in the mean wind speed and direction (Hurrell et al., 2003). In periods of higher than average pressure difference (positive NAO phase), when both the tropical high and polar low pressure features are relatively stronger, the north-south gradient produces an enhanced zonal (west to east) flow. When both the tropical high
and polar low pressure features are relatively weak (negative NAO phase), the synoptic scale flow has a reduced zonal component. Thus, influences on the interannual variability of hemispheric-scale pollution transport pathways related to the NAO are expected.

The study of the NAO impact on pollution transport is further motivated by results from several previous relevant analyses. Li et al. (2002) imply that the NAO can be used to forecast transatlantic transport of North American pollution to Europe based on a 5 yr simulation of tropospheric chemistry evaluating North Atlantic coastal station measurements. They find that the strength of the prevailing westerlies favors transport close to the surface. Creilson et al. (2003) also find that the NAO aids the transport of anthropogenic pollution from North America to Europe by influencing the westerly winds across the North Atlantic. Eckhardt et al. (2003) have studied the control that the NAO exerts over pathways of pollution transport towards the Arctic simulating fixed lifetime passive tracers over 15 yr and found that there is a strong relation between the NAO and surface concentrations of carbon monoxide (CO) in the Arctic, most pronounced during winter and spring. They also note that European pollution can build-up in the Arctic, with faster poleward transport during the positive phase of the NAO. In the present paper we study the emission and transport of tagged idealised tracers, with the characteristics of carbon monoxide (CO), and aerosols in the atmosphere, simulated over a 50 yr period, using the ECHAM/MESsy Atmospheric Chemistry (EMAC) climate model.

CO is a sensitive tracer for long-range transport of pollution as it is directly emitted by combustion sources and has a lifetime of 1–3 months in the atmosphere. Anthropogenic CO emissions are distributed similarly to those of many gaseous and particulate anthropogenic pollutants. To obtain a realistic description of tracer lifetime and abundance in the atmosphere we include a removal mechanism through oxidation by the hydroxyl (OH) radical, the main chemical species controlling the oxidizing capacity of the atmosphere. The OH radical is highly reactive and oxidises CO to form carbon dioxide CO$_2$. 

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Aerosols are a major component of atmospheric pollution and have important effects on human health (Huntingford et al., 2007), the environment and hydrological cycle (Ramanathan et al., 2001) and climate (Isaksen et al., 2009). To study the impact of the NAO on tropospheric aerosol pollution we simulated the emission and transport of idealised water-soluble aerosols. To decouple from the effects of initial distributions and allow for the direct comparison of the NAO effects on aerosol transport with those on trace gases, and to highlight differences in removal mechanisms, identical emission patterns are used for gases and aerosols. Aerosols in the present model setup are not chemically active but are affected by sedimentation and dry and wet deposition processes and are predominantly removed from the atmosphere by precipitation.

We briefly describe the model in Sect. 2, including the emissions of tagged tracers and removal mechanisms (Sect. 2.1). The simulation of the spatial structure and temporal evolution of the NAO is presented in Sect. 2.2. Seasonal averages for simulated fields are presented throughout the paper. The analysis of our results regarding the influence of the NAO on the intercontinental transport of tracer gases and aerosols is discussed in Sect. 3. The summary and conclusions are given in Sect. 4.

2 Model simulation

The EMAC model is a numerical chemistry and climate simulation system that includes sub-models describing tropospheric and middle atmosphere processes and their interaction with oceans, land and human influences (Jöckel et al., 2005). It uses the first version of the Modular Earth Submodel System (MESSy1) to link multi-institutional computer codes. The core atmospheric model is the 5th generation European Centre Hamburg general circulation model (ECHAM5, Roeckner et al., 2006).

For the present study we applied EMAC version 1.9 (based on ECHAM5 version 5.3) on the T42L19 resolution, i.e. with a spherical spectral truncation of T42 (corresponding to a quadratic gaussian grid of approximately 2.8 by 2.8 degrees in latitude and longitude) with 19 vertical hybrid pressure levels up to 10 hPa. The sea surface tem-
perature and sea ice distribution for AMIP II simulations (Taylor et al., 2000) between the years 1960–2010 were used as boundary conditions. The applied model setup comprised the submodels MECCA1 (Sander et al., 2005) for tropospheric and stratospheric chemistry, though highly simplified, RAD4ALL for radiation and heating processes, CLOUD for cloud cover and micro-physics processes including precipitation, and CONVECT for the vertical transports of trace species associated with convection. The DRYDEP (Kerkweg et al., 2006) and SCAV (Tost et al., 2006, 2007) submodels are used to simulate aerosol dry and wet deposition processes respectively.

Large-scale stratiform and small-scale convective precipitation are the dominant mechanisms for the removal of water-soluble chemically inert aerosols in the model. Our simulation is shown to reproduce the basic features of the average distribution of precipitation observed by Adler et al. (2003, Global Precipitation Climatology Project (GPCP)) in the region of interest between 1979–2004 in Fig. 1. In the domain 20°–80° N and 90° W–40° E, average precipitation in excess of 10 cm month⁻¹ follows a southwest to northeast pattern over the Atlantic Ocean with maximum values of the order of 15–20 cm month⁻¹ occurring off the eastern coast of the United States. Finer scale features over the southeastern United States and the western Scandinavian coast are also reproduced. The model overestimates precipitation over some parts of southern Europe close to the Mediterranean Sea though with an extend and magnitude that is expected to have limited influence on the results.

2.1 Emissions

The emission scheme of Pozzer et al. (2011) was adapted for the purposes of this study. Anthropogenic fossil fuel and biofuel CO emissions were simulated based on the global inventory prepared in the framework of the CIRCE project by the Emission Database for Global Atmospheric Research (EDGAR) of the EC-Joint Research Center Ispra, Climate Change Unit (Doering et al., 2009a,b). Emissions by international aviation were calculated using a technology based emission factor approach (Eyers et al., 2004). Production of CO from international shipping is based on the QUANTIFY project.
The vertical distribution of anthropogenic emissions, is based on experience from the European Monitoring and Evaluation Programme (EMEP) model (Simpson et al., 2003), and is described in Pozzer et al. (2009). Biogenic emissions have been compiled following Guenther et al. (1995) and are prescribed offline in the model. The contribution to emissions by biomass burning was included using the Global Fire Emissions Database (GFED version 3, van der Werf et al., 2010).

We use the base year 2005 with a monthly temporal resolution climatology for all emissions, except for biomass burning for which a 10 yr monthly average climatology (2000–2009) is used. The CO tracer is tagged by the regions of emission, i.e. from North America and Europe (see Fig. 2). Emissions from eastern Russia are very small and are not included in our definition of Europe. We do not account for the atmospheric chemical production of CO (≈ 1000 Tg yr⁻¹, Seinfeld and Pandis, 1997).

Loss of the tagged CO tracers is computed using the monthly atmospheric concentration of OH from the full-chemistry simulation in Pozzer et al. (2011). CO is removed from the atmosphere via the mechanism:

\[
\text{CO} + \text{OH} \rightarrow \text{CO}_2 + \text{HO}_2
\]

The small-scale recycling of OH radicals, e.g. by the reaction of NO with HO₂, and the large-scale buffering through ozone (Lelieveld et al., 2004) lead to the simplification:

\[
\text{OH} \Leftrightarrow \text{HO}_2
\]

in the model, and therefore we apply the reaction:

\[
\text{CO} + \text{OH} \rightarrow \text{CO}_2 + \text{OH}
\]

while keeping the atmospheric concentration of OH nudged towards monthly prescribed fields.

Aerosols are emitted using regional emission patterns and concentrations identical to those of CO. This facilitates the direct comparison of the effects of the NAO on the transport of trace gas and aerosol pollutants and their removal mechanisms. Water-soluble aerosols with a standard lognormal distribution of mean radius 0.5 µm (\(\sigma = 1.0\))
and a molar mass of 96.076 g mol\(^{-1}\) (equal to sulfate) with a Henry coefficient equal to 1.0 mol L\(^{-1}\) atm\(^{-1}\) and a density of 1841.0 kg m\(^{-3}\) are used.

### 2.2 North Atlantic Oscillation

There is no unique way to define the spatial structure and temporal evolution of the NAO. We applied several common approaches to evaluate our model. The NAO primary centers of action are identified in maps of 500 hPa geopotential height distributions as regions of maximum anti-correlation over the North Atlantic (Fig. 3). The 500 hPa height time series at a reference point (65° N, 30° W) is correlated with the corresponding time series at all points, revealing a north-south dipole, with one centre over the North Atlantic close to the Arctic and the other over the subtropical Atlantic, as expected.

The NAO signal is not readily apparent in the empirically determined orthogonal function (EOF) spatial pattern of the observed Northern Hemisphere (NH) winter 500 hPa height, but rather rotated (linearly combined) EOFs (Hurrell et al., 2003) have to be used. This is not an issue at the surface level. The spatial pattern reflected in the leading EOF of the North Atlantic sector (20°–80° N, 90° W–40° E) seasonal (3-month average) sea level pressure (SLP) anomalies field (Fig. 4, top) explains 38% of the total variance and compares well with the NAO pattern observed by Hurrell (2010). In this approach, the NAO is identified from the spatially and temporally mutually orthogonal eigenvectors of the cross-covariance matrix of time varying values of SLP. Again, a north-south pressure difference dipole over the Atlantic Ocean in the northern hemisphere is apparent. By regressing the SLP anomalies on the principal component time series the pattern is obtained in terms of pressure amplitude, with maximum departure of the field at the centre of the poles of the order of ±3 hPa, in good agreement with the equivalent analysis of observations by Hurrell (2010).

The normalised principal component time series of the leading EOF is correlated to the model seasonal NAO index (determined as the difference in normalised sea level
pressure between a stationary point at Ponta Delgada, Azores, and a stationary point in Stykkishólmur/Reykjavík, Iceland; adapted from Hurrell, 1995) with a correlation coefficient of 0.93 (Fig. 4, bottom). This result, similar to observations, further indicates that the leading EOF captures the temporal variation also described by the NAO index and allows to interchange the use of the pressure difference index and the EOF time series to diagnose the temporal variation of the pattern.

3 Results

3.1 Vertically integrated transports

The vertically integrated tracer transport vector is defined as

$$Q = \frac{1}{g} \int_{0}^{p_S} q u dp$$

where $q$ is the tracer concentration in mol mol$^{-1}$, $u$ the horizontal vector wind, and $p$ is pressure; $p_S$ denotes surface pressure and $g$ represents gravity. This variable is analogous to the vertically integrated moisture transport (see Fig. 5), which was used to test our model simulation against observations compiled by Hurrell (1995). During seasons of low or negative NAO index moisture transport over the Atlantic Ocean follows a westerly zonal path with maximum vector magnitudes occurring in the western part of the domain, stretching across the mid-Atlantic. During seasons of high NAO index the axis of maximum moisture transport shifts to a more southwest-to-northeast orientation over the Atlantic and extends farther to the northern and eastern part of Europe.

We calculated the temporal average of $Q$ for all winters with high (>1) and with low or negative (<0) EOF principal component time series values. In total, 10 high and 23 low or negative EOF time series seasonal averages enter the composites. The axis of maximum westerly North American CO tracer transport extends much farther
to the north and east, over northwestern Europe and Scandinavia, during seasonal composites with high NAO phase when compared to seasonal composites with low or negative NAO phase. This is indicated by vector plots of the vertically integrated total tracer transports (Fig. 6, top). The equivalent analysis for the CO tracer originating in Europe shows similar results over the Atlantic Ocean and a relative increase in transport over mainland Europe towards the East (Fig. 6, bottom).

### 3.2 NAO index – tracer concentration correlation

After identifying the principal component time variation, we examine the relation of the NAO with tracer concentrations. The correlation of the principal component time series with gas and aerosol concentration anomalies has been calculated for the model surface level and for the meridional vertical profile at 30° W. The results can be seen in Figs. 7 and 8 for the gaseous tracers and aerosols originating from the North American and European regions, respectively.

The principal component time series, and hence the phase of the NAO, is significantly correlated to North American CO tracer concentration anomalies over the northwestern Atlantic Ocean, and is anti-correlated to CO tracer concentrations over the tropical Atlantic Ocean (Fig. 7) close to the surface. For soluble aerosol species originating from North America, the effect is stronger and reversed with higher significantly positive correlations extending much farther over the North Atlantic across northwestern Europe. The patterns extend vertically, spanning the free troposphere with strong correlations around 60° N and anti-correlations at sub-tropical latitudes, more pronounced for the case of soluble species. The anti-correlation for aerosols extends to the equator and even over Africa, probably an effect of the increased moisture and precipitation over the tropics for seasons with low NAO index.

Similarly, the phase of the NAO is significantly correlated to European tracer concentration anomalies at the surface over high latitudes, within the Arctic circle, and over the Middle East, and anti-correlated over the Eastern Atlantic Ocean (Fig. 8). Contrary to the emissions from North America, for those from Europe the correlation patterns
for gaseous and aerosol tracers are not reversed. Similarly to the pollution originating in North America, the effect is stronger for the soluble species, extending over North Africa and mainland Europe. The anti-correlation over Northern Europe for the water-soluble aerosols may be explained by the effect of the NAO on the spatial distribution of precipitation, the main removal mechanism in the model. Swings in the NAO index are reflected in the transport and convergence of atmospheric moisture. High NAO index winters exhibit drier conditions over much of central and southern Europe, the Mediterranean and parts of the Middle East, whereas more than average precipitation falls over Scandinavia and Iceland (Hurrell, 1995). In the vertical profiles, a strong anti-correlation centred at 500 hPa, with a vertical extent from the surface to the tropopause region is most apparent for gaseous tracers, along with a weak surface correlation both at high and low latitudes. Interestingly, the gaseous tracer anomalies are highly anti-correlated with the NAO in the lowermost stratosphere at ∼200 hPa, which could be further investigated with an EMAC model version of the middle atmosphere (Jöckel et al., 2006). There are marked differences with the vertical profile for water-soluble aerosols, with a relatively strong correlation at tropical latitudes, which we attribute to the influence of the NAO phase and amplitude on the locations of high moisture and precipitation.

4 Conclusions

A 50 yr tracer transport simulation with the EMAC model was used to study the influence of the NAO on the distribution of atmospheric pollution. Regionally tagged tracers of carbon monoxide and water-soluble aerosols were emitted using a fixed emission climatology, taking into account anthropogenic (fossil fuel, biofuel, aviation and shipping) sources, natural sources and biomass burning. Removal mechanisms were included based on oxidation of CO by the OH radical and sedimentation and dry and wet deposition of aerosols. The model simulation of the boreal winter NAO spatial pattern was tested against observations using common and established approaches. Good
agreement with results reported from observations was obtained throughout.

The NAO phase is related to pollution transport from North America towards Europe, mostly in the free troposphere. The NAO is associated with stronger-than-average westerlies across the middle latitudes of the Atlantic that transport air pollution from North America towards Europe and Northwestern Africa, similar to moisture transport.

The NAO exerts a clear influence on the atmospheric distribution of aerosol pollution. The NAO phase strongly affects the magnitude and direction of moisture transport, including water-soluble species emitted in North America and Europe, over the Atlantic Ocean. The NAO also affects the location and extent of the spatial pattern of precipitation. During high NAO index periods, drier than average conditions occur over much of central and southern Europe, the Mediterranean and parts of the Middle East, whereas more precipitation than average falls over a region from Iceland through Scandinavia (Hurrell, 1995). As precipitation is the main mode of removal of water-soluble aerosols from the atmosphere, the distribution of particulate pollutants, especially from Europe, is indirectly influenced by the NAO. The response of the aerosols is relatively strong, as they are influenced both by NAO-related transport and wet deposition anomalies, whereas CO is unaffected by precipitation.

Hurrell (1995) concludes that decadal variability in the NAO has become especially pronounced since 1950, with unprecedented strongly positive index values since 1980. If this trend persists, e.g. associated with climate change (Intergovernmental Panel on Climate Change, 2007), it can be expected that in the future the NAO will have a growing influence on intercontinental pollution transport between North America and Europe and from Europe towards the Arctic.

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Fig. 1. Observed (Adler et al., 2003, Global Precipitation Climatology Project (GPCP)) (left) and modeled (right) average precipitation (in cm/month) between 1979–2004.
Fig. 2. Depiction of the regions used for the computation of emissions.
Fig. 3. One-point correlation map of model 500 hPa geopotential height for boreal winter (December-February) over the period 1960–2010. The reference point is 65° N, 30° W.
Fig. 4. Top: Leading empirical orthogonal function (EOF1) of the winter (December-February) mean sea level pressure (SLP) anomalies in the North Atlantic sector (20°–80° N, 90° W–40° E) which explains 38% of the total variance. The pattern is displayed in terms of amplitude (hPa), obtained by regressing the SLP anomalies on the principal component time series. Bottom: Principal component time series (red, blue) of the leading EOF plotted with the modeled NAO index (black) based on the normalised SLP difference between Ponta Delgada, Azores, and Stykkishólmur/Reykjavík, Iceland. The correlation coefficient is 0.93.
Fig. 5. Average vertically integrated moisture (in kg/kg) transport vectors for winters with low or negative (left) and high (right) EOF time series. Red contour lines correspond to vector magnitudes.
Fig. 6. Average vertically integrated CO tracer (in mol mol$^{-1}$) transport vectors with North American origin (top) and European origin (bottom) for winters with low or negative (left) and high (right) EOF time series. Contour lines correspond to vector magnitudes.
Fig. 7. Correlation of winter (December-February) seasonal surface concentration anomalies (top) and meridional vertical profiles at 30° W (bottom) with the principal component of the EOF time series of SLP in the North Atlantic sector for gaseous tracers (left) and soluble aerosols (right) emitted from North America. White dashed contour lines indicate the local 95 % level of significance.
Fig. 8. Correlation of winter (December-February) seasonal surface concentration anomalies (top) and meridional vertical profiles at 30° W (bottom) with the principal component of the EOF time series of SLP in the North Atlantic sector for gaseous tracers (left) and soluble aerosols (right) emitted from Europe. White dashed contour lines indicate the local 95% level of significance.