Atmospheric Trajectory and Chemical Transport Modelling for Elevated Ozone Events in Denmark

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ABSTRACT

In this study, three Danish sites having the longest (1990-2004) time-series of ozone measurements were analysed on inter-annual, monthly and diurnal cycle variability as well as elevated and lowered ozone concentration events were identified. The atmospheric trajectory (HYSPLIT) and dispersion (HIRLAM + CAMx) models were employed to study dominating atmospheric transport patterns associated with elevated events and to evaluate spatio-temporal variability of ozone specific episode and typical seasonal patterns for Denmark. It was found that generally inter-annual variability has a positive trend, and events with low ozone concentration (≤10 μg/m³) continued to diminish. On a monthly scale, the highest and lowest mean concentrations are observed in May and November-December, respectively. The elevated concentrations (≥120 μg/m³) are observed during March-September. On a diurnal cycle, it is observed mostly during 13-16 of local time, and more frequent (ten-fold) compared with nighttime-early morning hours. For ozone elevated events, several sectors (or pathways of atmospheric transport) were identified depending on the sites’ positions, showing the largest (39%) number of such events associated with the north-western sector, and lowest (13% each)–southwestern and northern sectors. For each site, less than 60 events showed very high concentrations (≥180 µg/m³). Among 12 episodes, one longest elevated episode (19-21 Jun 2000) simultaneously registered at all sites and characterized by dominating transport from the south-southwestern sector, low wind speed, clear-sky, and multiple inversions was studied using modelling tools. For this episode, both measurements and modelling (trajectory and dispersion) results showed a relatively good agreement.

Keywords: Elevated Ozone Concentration Event and Episode; Atmospheric Trajectory; Chemical Transport Modeling; HYSPLIT; HIRLAM; CAMx

1. Introduction

Ozone (O₃) is one of the pollutants of the major concern for population and environment, and its high levels are an indicator of high pollution representing significant risks to humans, flora, fauna [1], as well as economic damage [2]. For several decades ozone has been operationally forecasted by many countries. It has been analyzed to get insight on local and regional trends, daily, seasonal and inter-annual variability, spatial and temporal distribution. Ozone is a natural component of the troposphere, produced by photochemical reactions of nitrogen oxides (NOₓ = NO + NO₂) and volatile organic compounds (VOCs) emitted from local and remote anthropogenic and biogenic sources. Once it has been produced it could exist in the troposphere for several days. Close to anthropogenic sources such as industrial or highly urbanized areas, emitted NO may react to form NO₂, and hence, to reduce ozone locally. Therefore, generally maximum concentration and exposure to ozone pollution will occur downwind of the sources [3], e.g. in suburban and rural areas. However, for some regions with specific topography and under certain synoptical conditions air masses can re-circulate and stay within a region for longer period of several days.

The major mechanisms responsible for elevated ozone during heat-wave episode included horizontal transport from continental Europe with existence of a long-lived anticyclone, convergence of westerly and easterly near-surface winds, or downward entrainment of ozone-rich air from residual layers in the free troposphere [4]. Ozone production highly depends on meteorological conditions including sunlight, cloudiness, temperature, and wind characteristics [5]. Ozone measured at specific site may be produced many hundred kilometres away.
can be transported from elsewhere in the world because intercontinental ozone transport is an efficient process [6]. Even application of various policy options for ozone precursors may still remain a significant potential for photochemical ozone formation and long-range transport [7]. Episodes of high ozone concentrations are the most interest; and these are based on a sum of existing regional background ozone levels, stratospheric ozone injections, and contribution of ozone produced in troposphere from naturally occurring and man-made precursors.

In Denmark, automatic ozone measurements are performed by the Danish National Environment Research Institute (NERI) in selected urban and rural areas. Such data are publically available at the European Monitoring and Evaluation Programme (EMEP; http://www.emep.int) website. In general, instruments record the regional background level for Denmark, but also, on occasion, pollution transported from sources outside of the country.

In this study, the long-term (1990-2004) time-series of ozone measurements at 3 sites have been analysed on inter-annual, monthly and diurnal cycle variability as well as cases with elevated and lowered concentrations were identified. Atmospheric trajectory and dispersion modelling tools were applied to study dominating atmospheric transport patterns associated with elevated ozone events and spatial and temporal variability of concentration patterns for selected episodes and on a seasonal scale for Denmark [8,9].

### 2. Methodology

#### 2.1. Ozone Measurements

The available original time-series of ozone measurements at three selected Danish sites—DK31 (56.28˚N, 08.43˚E; Ulborg), DK32 (55.97˚N, 12.33˚E; Frederiksvborg), and DK41 (55.69˚N, 12.13˚E; Lille Valby) covered a period from 1990 to 2004. Two of these sites (DK41 and DK32) are located on the Zealand island and one site (DK31) is placed on the Jutland Peninsula (Figure 1(a)) of Denmark. After pre-screening and removing all missing and below detection limit values, the total number of available measurements for analysis had been reduced (Table 1). Each record included information on a value of concentration and corresponding temporal parameters (year, month, day, hour of measurement).

General statistics from analysis of the revised time-series showing a number of the cases with different elevated (i.e. above 120, 150, and 180 μg/m³) and lowered ozone concentrations (i.e. below 10 μg/m³) as well as years with highest and lowest number of such cases in selected years for three sites is summarized in Table 1. As seen, during the period studied the absolute maximum value of concentration was above 200 μg/m³ for all 3 sites. The elevated ozone levels were registered only in about 1.5% and less than 1% of the cases for the DK31 and DK41, DK32 sites, respectively. Note that ozone levels above 120 μg/m³ were registered practically in all years analyzed, but very high levels (above 180 μg/m³) were observed only in 5 years from all 15 years considered. Note that observed peaks in ozone concentration

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**Figure 1.** (a) Geographical positions (extracted from Google-Earth) of Danish sites for ozone measurements (A: DK31, DK32, and DK41), synoptical meteorological stations (W: WMO-06160—Vaerlose, WMO-06168—Nakkehoved Fyr, WMO-06058—Hvide Sande, WMO-06052—Thyboron) and sounding stations (S: Jaegersborg, Karup); and (b) Boundaries of domains for atmospheric (NWP_G45) and chemical transport (ACT-CAMx) modelling.
Table 1. Summary statistics on a number of the cases (N) with ozone concentration levels (above 120, 150, and 180, and below 10 $\mu$g/m$^3$) for three Danish measurement sites (H—highest, L—lowest).

<table>
<thead>
<tr>
<th>Measurement Site Parameter</th>
<th>DK-31</th>
<th>DK-41</th>
<th>DK-32</th>
</tr>
</thead>
<tbody>
<tr>
<td>Original # measurements</td>
<td>131495</td>
<td>122735</td>
<td>105192</td>
</tr>
<tr>
<td>Revised # measurements</td>
<td>115690 (100%)</td>
<td>115447 (100%)</td>
<td>97939 (100%)</td>
</tr>
<tr>
<td>N (O$_3$ &gt; 120 $\mu$g/m$^3$)</td>
<td>1745 (1.5%)</td>
<td>1079 (0.9%)</td>
<td>811 (0.8%)</td>
</tr>
<tr>
<td>N (O$_3$ &gt; 150 $\mu$g/m$^3$)</td>
<td>206</td>
<td>161</td>
<td>140</td>
</tr>
<tr>
<td>N (O$_3$ &gt; 180 $\mu$g/m$^3$)</td>
<td>36</td>
<td>42</td>
<td>31</td>
</tr>
<tr>
<td>N (O$_3$ &lt; 10 $\mu$g/m$^3$)</td>
<td>2990 (2.6%)</td>
<td>8732 (7.6%)</td>
<td>8464 (8.6%)</td>
</tr>
<tr>
<td>Abs Max O$_3$, $\mu$g/m$^3$</td>
<td>221 (1990)</td>
<td>213 (1992)</td>
<td>213 (1992)</td>
</tr>
</tbody>
</table>

are dependent on regional background levels of ozone, specific rate of photochemical production during a day, and contribution of other chemical species (such as nitric oxide and VOCs) to ozone related chemical reactions. In addition, both local and remote sources and on occasion, stratospheric ozone injections could play an important role. The lowered ozone levels were observed more frequently (up to 8.6%), and a number of such cases was larger for sites of the Zealand compared with Jutland regions.

2.2. Atmospheric Trajectory Modelling

The backward trajectory modelling is a widely used tool for evaluation of possible atmospheric transport to geographical locations where short- or long-term measurements of chemical species are conducted. The tracking of air parcels spatial positions in the atmosphere during such transport can allow identifying potential paths and regions from where pollution can be transported and associated. The accuracy of the trajectories is typically of the order of 20% of the travelling distance, although in some cases it can be larger [10].

In our study, the National Oceanic and Atmospheric Administration (NOAA) on-line transport and dispersion Hybrid Single-Particle Lagrangian Integrated Trajectory model (HYSPLIT) v 4.5 model available in an interactive mode http://www.arl.noaa.gov/ready/open/hysplit4.html was used [11,12]. HYSPLIT performs computing of both forward and backward types of trajectories using several methods. To drive trajectory simulations the gridded meteorological dataset (http://dss.ucar.edu/pub/reanalysis)—NCEP/NCAR Global Reanalysis 1948-Present—was used [13]. It has temporal coverage 4 times per day (00, 06, 12, 18 UTCs), including daily and monthly values from 1 Jan 1948 to present. Spatial coverage includes entire globe. There are 17 pressure level and 28 sigma levels.

Each computed trajectory was associated with the corresponding elevated ozone measurement at one of the three sites. For simplicity, only one trajectory arriving at the measurement sites at the ground level for term with the highest value of concentration was computed backward in time up to 120 hours (5 days) using vertical motion calculation method. This duration was selected because of the residence time of ozone in the atmosphere which is leading to a characteristic scale of its atmospheric transport of 1000 km. A set of meteorological data was also extracted along each trajectory at 6 hour intervals. It included the terrain height, potential temperature, air temperature, precipitation, relative humidity, and mixing layer height. Because there is a difference of 1 hour between the universal coordinated time (UTC) and Danish LST, the trajectories were computed at corrected times in order to fit the corresponding measurements done at local times. Then all trajectories were grouped depending on pathway of atmospheric transport and area from where trajectories have arrived.

2.3. Meteorological and Chemical Transport Modelling

2.3.1. Meteorological Modelling

The Numerical Weather Prediction (NWP) models (with their 3D meteorological output) are used as drivers for the Atmospheric Chemistry Transport (ACT) models. The selection of meteorological driver as well as boundaries for domain of interest will depend of specific application. Hence, different meteorological operational or reanalyzed archived datasets from NWP can be used. Output from operational and climatological models can be
used in different applications.

Danish Meteorological Institute (DMI) runs several nested versions of the HIRLAM (http://hirlam.org) model with horizontal resolution ranging from 15 to 3 km and forecast length up to 72 hours. Moreover, there is an urbanized version of HIRLAM model (resolution 1.5 km) run in a research mode for environmental applications. The operational DMI forecasting modelling system [14,15] includes the pre-processing, climate file generation, data assimilation, initialization, forecasting, post-processing, and verification. It includes also a digital filtering initialization, semi-Lagrangian advection scheme, and a set of physical parameterizations such as Savijærvi radiation [16], STRACO condensation [15], CBR turbulence scheme [17], and ISBA land surface scheme [18]. The lateral boundary conditions are received every 6 hour from European Center for Medium-Range Weather Forecasts (ECMWF). The system is running on the DMI CRAY-XT5 supercomputer and produced output is archived on a mass storage system. In our study, the DMI-HIRLAM-G45 model (45 × 45 km, 31 vertical levels) was run to produce 3D meteorological fields (see domain in Figure 1(b)).

2.3.2. Chemical Transport Modelling

The chemical transport modelling is based originally on the CAMx (Comprehensive Air quality Model with extensions) model (http://www.camx.com) with built an interface between NWP (DMI-HIRLAM-G45) and ACT (modified CAMx, v 4.4) models. Through interface necessary information was extracted from the NWP model output, and then it served as input for the ACT model. The Carbon Bond IV (CB-IV) Mechanism [19] with is used. This mechanism is used together with the Tropospheric Ultraviolet and Visible (TUV) radiation model [20] to calculate photolysis rate coefficients. The modelling system is used to simulate aerosols and gas-phase compounds from different scales of ground-level gas-phase air pollutants. It can be used for air quality forecasts and modelling of historical data.

The horizontal and vertical resolutions of the ACT model depend on a resolution of the used meteorological and emission data. The selected ACT-CAMx domain (Figure 1(b)) includes the European countries with Denmark in the center. It has a horizontal resolution of 0.2° × 0.2° (apprx. 20 × 20 km) and 25 vertical levels including 15 HIRLAM levels covering the lowest 3 km of the troposphere. The amount of chemical compounds, which is transported from the free troposphere into the atmospheric boundary layer, is determined by the meteorological information and concentration of chemical compounds in the free troposphere. These concentrations depend on the latitude, longitude, land/sea mask and month of the year. The TNO MEGAPOLI emission inventory [21] (0.06° × 0.12° resolution) had been interpolated into grids of the ACT model taking into account vertical distribution. In order to reach a chemical equilibrium, a 2 day spin up has been applied, and the forecast length extended up to 24 hours.

Moreover, in order to study the chemical transport and links with dominating meteorological conditions, it is necessary to analyze in more details the available surface maps, vertical sounding diagrams, and observations at synoptical meteorological stations (Figure 1(a)) located near the ozone measurement sites.

3. Results and Discussions

3.1. Observed Inter-Annual, Monthly and Diurnal Variability

Analysis of inter-annual variability (during the period studied 1990-2004) of ozone time-series showed generally a positive trend of ozone increase (Figure 2(a)). The number of cases with lowered ozone concentration levels (i.e. below 10 μg/m³, see Figure 2(b)) continued to decrease throughout the period of measurements, and it occurs more rapidly for sites located on the Zealand Island (DK32 and DK41) compared with DK-31 site on the Jutland Peninsula. In general, for a period of more than 10 years, the number of such cases decreased almost twice.

Month-to-month variability of the mean ozone concentration show clear differences between the sites (Figure 2(c)). As seen, throughout the year, the higher mean ozone levels are observed for the DK31 site, compared with others; and the lower—for DK32 site. The highest mean concentrations are characteristic in May for all sites, and for DK31 site it is the largest of about 80 μg/m³. The mean concentrations are the lowest in November-December, decreasing to about 40 and 30 μg/m³ for the DK31 and DK32, DK41 sites, respectively. The elevated concentrations (i.e. more than 120 μg/m³) are observed during March-September (Figure 2(d)). Throughout the mentioned period the number of such cases is larger for the DK31 site compared with two others. The maximum number of such cases is in August for DK31 and DK41 sites, compared with DK32—in June.

On a diurnal cycle, all sites showed similar shape of variability with a maximum occurred at about midday (14 hour of the local time) and a minimum—in the morning (at 06 hour) (Figure 2(e)). A shape of such diurnal variability is due to high reactivity of ozone. Concentration at the surface level is decreased rapidly in evening hours (although it could remain elevated above the mixing layer and then contribute to elevated ozone levels in the following days). In time-series, the absolute ozone maxima (considering levels of above 120 μg/m³) are observed mostly during 13 - 16 hours, and the number of such cases is ten-fold larger compared with nighttime-
Figure 2. Ozone inter-annual ((a) and (b)), monthly ((c) and (d)) and diurnal cycle ((e)-(h)) variability for the mean concentration in μg/m³ ((a), (c) and (e)), absolute maximum concentration (f), number of cases with concentrations below 10 μg/m³ ((b) and (h)) and above 120 μg/m³ ((d) and (g)) for three Danish sites DK31 (--), DK32 (---) and DK41 (----).
early morning hours (Figures 2(f) and (g)). The number of the cases with the lowest ozone concentration (less than 10 μg/m³) varied from about 100 (DK31) to 200 (DK32) cases during 11 - 15 hours, and it was up to 3 times higher during late evening-nighttime hours for the Zealand sites, but almost non-variable for the Jutland site (Figure 2(h)).

Note that on figures for the mean ozone concentration (i.e. Figures 2(a), (c) and (e)) the similar pattern of higher ozone concentrations on inter-annual, monthly, and diurnal cycle scales exists for the DK31 site compared with two other locations.

3.2. Atmospheric Transport Sectors and Pathways

Distribution of trajectories (number and percentage) arrived at three measurement sites during 1990-2004 by sectors and pathways, linked with atmospheric transport, is summarized in Table 2. As seen, the clustering of trajectories revealed 7 possible atmospheric transport pathways divided into sectors (i.e. from where trajectories have arrived). In total, 207, 139, and 160 trajectories were associated with elevated ozone events observed at the DK31, DK32, and DK42 measurement sites, respectively.

The largest number of trajectories connected with such events was associated with the North-West sector (in total 155 trajectories). Trajectories from this sector are all originated over the Atlantic Ocean aquatoria. But there are several paths of such transport through: 1) UK, Benelux countries, and Germany; 2) UK and North Sea (only for DK31 site; 9 trajectories); 3) Norway and North Sea; and 4) Benelux countries and Germany. The first path dominates among others (more than 20 trajectories per each site). The smallest number of trajectories was associated with the South-West sector (where trajectories also originated over Atlantic Ocean, but additionally passed over France and Benelux countries before reaching Denmark) for DK31 and DK32 sites, and with the North sector—for DK41 site.

Trajectories, arriving at the three measurement sites from the North sector, are all originated over Scandinavian countries (Norway and Sweden). There are two possible comparable paths of such transport through: 1) Baltic Sea and Poland/Germany (28 trajectories); and 2) Baltic Sea (26 trajectories). Note, that the first path is more characteristic for the DK31 site (18 trajectories), but the second path is almost equally (about 8 trajectories per site) observed at all three sites.

Atmospheric transport from the South sector (through Germany/Benelux countries) is the second largest (accounting for 81 cases). Moreover, some trajectories did not show clear pathway of atmospheric transport (i.e. had complex structure and circulated over Denmark) and these were assigned to “no clear identification” category. In total, this category includes 34 trajectories (out of 506).

Several examples of simulated backward trajectories associated with elevated ozone arriving at sites underline dominating direction of atmospheric transport of air parcels toward the measurement sites’ locations (as seen in Figure 3). In figures a distance between marked 12 h intervals shows approximate speed of such transport. The changing spatial position of trajectory shows movement of air parcels through possible source regions. For example, as seen in Figure 3(a), air parcels arriving at the DK41 site, located on the Zealand Island, originated from the Atlantic Ocean aquatoria 5 days before. These have traveled through the UK metropolitan and industrial areas as well as the North Sea on its way to Denmark. This pathway is associated with atmospheric transport from the South-West sector. Figure 3(b) depicts that air parcels arrived at the DK31 site, located on the Jutland Peninsula, originated over Benelux countries and heavily populated and industrialized Rein-Ruhr area (Germany). This example is associated with atmospheric transport from the South sector. Figure 3(c) presents transport from the East sector. In particular, the trajectories originated over the North-West Russia passed through the Baltic States and aquatoria of the Baltic Sea before reaching the DK31 site.

Table 2. Summary (number and percentage) of trajectories arrived at 3 ozone measurement sites during 1990-2004 by sectors and pathways of atmospheric transport.

<table>
<thead>
<tr>
<th>Sector</th>
<th>N traj by sector</th>
<th>Pathway</th>
<th>% of trajectories by site</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>DK31</td>
</tr>
<tr>
<td>E</td>
<td>71</td>
<td>BST (RU) → Sea → PL → (GE)</td>
<td>12.1</td>
</tr>
<tr>
<td>SW</td>
<td>43</td>
<td>AO → FR → Blux/GE</td>
<td>5.8</td>
</tr>
<tr>
<td>S</td>
<td>81</td>
<td>GE/Blux</td>
<td>16.8</td>
</tr>
<tr>
<td>SE</td>
<td>68</td>
<td>PL/GE</td>
<td>6.8</td>
</tr>
<tr>
<td>NW</td>
<td>155</td>
<td>From AO</td>
<td>38.7</td>
</tr>
<tr>
<td>N</td>
<td>54</td>
<td>From Scandinavia</td>
<td>13.1</td>
</tr>
<tr>
<td>NOID</td>
<td>34</td>
<td>No clear identification</td>
<td>6.7</td>
</tr>
<tr>
<td></td>
<td>506</td>
<td>total % of trajectories by site</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td></td>
<td>total number of trajectories</td>
<td>207</td>
</tr>
</tbody>
</table>

Comment: AO—Atlantic Ocean; BST—Baltic States; RU—Russia; PL—Poland; GE—Germany; FR—France; Blux—Benelux countries; Scand.—Scandinavian countries (Norway/Sweden).
3.3. Elevated Concentration Episodes

Several episodes with elevated levels were identified from detailed analysis of multi-year ozone time-series. For such episodes the level of concentration depends on both the local conditions and due to long-range transport contribution. The following criteria were chosen: 1) concentration higher 150 μg/m³; 2) observed, at least, during 24 hours period; 3) simultaneously, at least, at two measurement sites; and 4) associated with one of the potential source regions (i.e. according to identified sectors of atmospheric transport (see Ch. 3.2). There were 7, 6 and 5 relatively long-term episodes for the DK31, DK32, and DK41 sites, respectively. The ranking of these episodes is the following: 1) 19-21 Jun 2000 and 30 Jun-1 Jul 1992—occurred at 3 sites simultaneously; 2) 4 Aug 1990, 9-10 Aug 1992, 26-28 Jul 1994 and 5 May 1995—occurred at 2 sites simultaneously; and 3) 28 Jul 1995, 6-8 Jun 1996, 6 Aug 1999 and 16 May 2000—occurred at one of the sites. Although elevated concentrations were also observed in other months, but duration of episodes was very short. One episode with the longest duration (i.e. 19-21 Jun 2000) was selected for detailed analysis employing the HIRLAM + CAMx modeling system.

3.4. Case Study—Episode for 19-21 June 2000

During 19-21 Jun 2000 elevated ozone episode, in the Danish studied area the typical meteorological conditions near the surface were characterized by dominating winds (with low speeds up to 5 m/s) from the south-southwestern sector (associated with Benelux countries and Germany). Although wind conditions in this sector were similar to Danish but it had been warmer there. During the episode the air temperature varied 16˚C - 30˚C (sector) compared with 14˚C - 24˚C (Denmark). Most time it was cloud-free and become partly cloudy on the last day of the episode. The relative humidity ranged from 48% to 97% on Jutland and from 41% to 99% on Zealand. The twice daily sounding diagrams from the Jægersborg (on Zealand) and Karup (on Jutland) stations showed the presence of several inversions in the low layers (some originated at the surface and some at higher altitudes, and extended up to 850 hPa). Moreover, on a preceding day (i.e. 18 Jun 2000) to the episode the isothermal layers were also observed at 600 - 650 hPa levels. The wind patterns remained of the south-southwestern sector and dominated within the boundary layer up to 850 hPa (with speeds up 5 - 10 and 5 - 15 m/s for the Jægersborg and Karup stations, respectively).

The elevated ozone episode in June 2000 started at 12 UTC on 19 June 2000 (see Figure 4(a)), i.e. when ozone concentration became above 150 μg/m³ at DK31 site. Then, concentration rapidly increased at this site reaching more than 200 μg/m³ within a few hours. Due to atmospheric transport the plume with elevated ozone levels reached the DK32 site around 08 UTC on 20 Jun 2000, and a few hours later—the third site DK41. On this day from 10 UTC, the elevated concentrations were recorded at all three sites simultaneously. On the same day at about 16 UTC, the absolute maxima 194 and 207 μg/m³ were observed at DK32 and DK41 sites, respectively.
maxima were observed almost a day later after the maximum at the DK31 site. Then, the concentration gradually decreased at all three locations, and this episode has ended after 17 UTC on 21 Jun 2000 (i.e. when concentration become below 150 μg/m³).

As seen in Figures 4(b)-(d), the atmospheric transport toward three measurements sites have occurred from the South-Western sector, i.e. mostly from the industrial and urbanized areas of the Benelux countries and north-western parts of the Rein-Ruhr region. Before arrival in Denmark during 3 days the air masses travelled mostly within the atmospheric boundary layer and passed over these regions getting additional ozone load.

An example of the CAMx simulated chemical transport patterns (Figure 5) is given on one of the days of the selected episode (nighttime-noon on 19 June 2000) with output provided at every 12 hour interval.

At nighttime, the ozone concentration is below 30 μg/m³ over the most area of the European domain, including Denmark and it had started to increase during morning hours reaching more than 130 μg/m³ at noon in the northern territories of Germany and Jutland Peninsula of Denmark where the DK31 site is located. The time-series shows almost similar value of measurements (150 μg/m³), which underlines that the CAMx model showed relatively good performance, although some discrepancies can be attributed to a relatively low horizontal resolution of the model run (i.e. when peaks in concentration associated with specific location will be smoothed over grid-cell). For comparison of elevated ozone concentration episode, the typical monthly concentration patterns for June 2000 are shown in Figure 6(a) (see explanation in Ch. 3.5).

3.5. Modelled Monthly, Seasonal and Annual Patterns Authors and Affiliations

The simulated monthly-averaged typical atmospheric chemical transport pattern (characteristic for the June 2000) over the European domain is shown in Figure 6(a). As seen, during June the ozone concentration over Denmark varied, on average, between 50 μg/m³ (for northern territories of Denmark) and 60 μg/m³ (southern territories

![Figure 4](attachment:image.png)
Figure 5. Spatial distribution of ozone concentration (in μg/m³) field for 19 Jun 2000 ((a) and (b): at nighttime-noon; 00-12 UTC) (simulated by HIRLAM + CAMx).
of Denmark and border regions with Germany). The territories southerly of Denmark have higher ozone levels (above 60 $\mu$g/m$^3$). These are associated with the Rein-Ruhr industrialized region and London metropolitan areas which are both located in a relative proximity to Denmark; and hence, polluted air masses can be transboundary transported toward Denmark.

The simulated seasonally-averaged patterns over the European domain, with a focus on territory of Denmark, were also analyzed. During spring the averaged ozone concentration over Denmark varied between 40 - 50 $\mu$g/m$^3$. The large area of the inland industrialized areas of France, Benelux countries, Germany, Po Valley (Italy) is associated with concentration higher than 50 $\mu$g/m$^3$. The concentration less than 40 $\mu$g/m$^3$ is observed over the large parts of the Scandinavia, Baltic States, Belorussia, and North-West Russia. During summer (Figure 6(b)), the situation is changed. In Denmark, the concentration has increased above 50 $\mu$g/m$^3$ for Zealand, Fyn and southern Jutland, except the northern and central parts of the Jutland Peninsula. The total area with the ozone level of 50 $\mu$g/m$^3$ expanded by almost a factor of 2, and it is more extended in the longitudinal compared latitudinal direction. The Rein-Ruhr area become more visible (above 60 $\mu$g/m$^3$) as well as the London and Paris metropolitan areas compared with others. The lower ozone (below 30 $\mu$g/m$^3$) is observed over the central and northern territories of the Scandinavia, and partially over the Baltic States and North-West Russia.

During fall, for the entire territory of Denmark the ozone level is slightly above 40 $\mu$g/m$^3$. Ozone is about 40 $\mu$g/m$^3$ over territories southerly of Denmark and within the 0 - 15°E longitudinal belt. The area with ozone level below 30 $\mu$g/m$^3$ moved far northward compared with summer; and it covers only central and northern territories of Sweden and Finland as well as Karelia and Kola Peninsula of Russia. As seen in Figure 6(c), during winter the ozone concentration is the lowest (below 30 $\mu$g/m$^3$) compared with other seasons analyzed. Such levels are also characteristics for the territories of European countries situated northerly 50°N.

On an annual scale, the averaged ozone concentration over Denmark varied between 40 - 50 $\mu$g/m$^3$. The lowest concentration is observed for the central territories (around 65 ± 2°N) of Scandinavia. For inland territories of the majority of the European countries (except the Baltic States, Belorussia, and Russia) the ozone levels are above 40 $\mu$g/m$^3$. The Rein-Ruhr and Po Valley Industrial
areas have higher ozone levels of about 50 \mu g/m^3.

4. Conclusions

In this study, three Danish measurement sites having the longest time-series of ozone measurements (with a time resolution of 1 hour and started in early 1990s) were evaluated—Ulfborg (DK31); and Frederiksborg (DK32) and Lille Valby (DK41). These sites are located on the Jutland Peninsula and Zealand Island of Denmark. After pre-screening of the time series covering almost 15 year period, it has been analysed on inter-annual, monthly and diurnal cycle scales with more detailed evaluation of cases characterized by elevated and lowered ozone concentrations. Both the atmospheric Lagrangian trajectory (HYSPLIT) and Eulerian dispersion (HIRLAM + CAMx) models were employed to study dominating atmospheric transport patterns (attributed to sectors) associated with elevated ozone events and to evaluate spatio-temporal variability of ozone concentration patterns for selected elevated episode in Jun 2000 and comparison with monthly (Jun) and seasonal patterns for Denmark.

It was found that ozone inter-annual variability showed generally a positive trend, and the number of cases with low ozone concentration (less than 10 \mu g/m^3) continued to diminish, and in particular, is has increased almost twice. Monthly ozone variability showed clear differences between measurements on Jutland vs Zealand. The highest mean concentrations are observed in May for all sites. The mean concentrations are the lowest in November-December. The elevated concentrations (more than 120 \mu g/m^3) are observed during March-September, and number of such cases is larger for Jutland (with maximum in August). On a diurnal cycle, ozone above 120 \mu g/m^3 is observed mostly during 13-16 of local time, and the number of such cases is ten-fold larger compared with nighttime-early morning hours.

For all ozone elevated cases (in total, 506), using trajectory analysis, several sectors or pathways of atmospheric transport were identified. Depending on the geographical position of the site, up to 39% of elevated events can be associated with transport from the north-western sector, from southern and south-eastern—22% each, eastern—18%, and from south-western and northern sectors—13% each. Among trajectories associated with elevated ozone levels, most of them travelled through inland (i.e. through the Baltic States, Russia, Poland, Germany, France, Benelux and Scandinavian countries) as well as over water areas (i.e. transport from the Atlantic Ocean, Baltic and North Seas).

Among ozone elevated events, less than 60 events for each site showed very high ozone concentrations (i.e. above 180 \mu g/m^3). Several long-term episodes with continuous elevated ozone levels were identified: 7 (in Jun 1996 and Jun 2000) and 5 (in Jul 1992 and Jun 2000) for the for Jutland and Zealand sites, respectively. One episode (19-21 Jun 2000) simultaneously registered at all three sites was studied in more details using the chemical transport model. This episode was characterized by low wind speed with atmospheric transport from the southwestern sector (associated with Benelux and German urban areas), mostly cloud-free conditions, and multiple inversions within the boundary layer. Measurements and modeling (trajectory and dispersion) results showed a relatively good agreement, although some discrepancies in concentration could be attributed to a relatively low horizontal resolution of the model run. For comparison, at the same time the typical monthly (June) ozone concentration varied between 50 - 60 \mu g/m^3 for northern vs. southern territories of Denmark. In spring and summer, the averaged ozone concentration varied 40 - 50 and above 50 \mu g/m^3, respectively. In fall, it is slightly more than 40 \mu g/m^3, and in winter—it is the lowest (less than 30 \mu g/m^3) throughout the year.

The results of this study are applicable for evaluation of short- and long-term ozone effects on health of population in both urban and rural areas and impact on environment (agriculture, forest, etc. damage), estimation of exposure levels for elevated episodes, and decision and policy making

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