Springtime nitrogen oxides and tropospheric ozone in Svalbard: results from the measurement station network

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Abstract. Svalbard is a remote and scarcely populated Arctic archipelago and is considered to be mostly influenced by long-range-transported air pollution. However, there are also local emission sources such as coal and diesel power plants, snowmobiles and ships, but their influence on the background concentrations of trace gases has not been thoroughly assessed. This study is based on data of tropospheric ozone (O\textsubscript{3}) and nitrogen oxides (NO\textsubscript{x}) collected in three main Svalbard settlements in spring 2017. In addition to these ground-based observations and radiosonde and O\textsubscript{3} sondes soundings, ERA5 reanalysis and BrO satellite data have been applied in order to distinguish the impact of local and synoptic-scale conditions on the NO\textsubscript{x} and O\textsubscript{3} chemistry. The measurement campaign was divided into several sub-periods based on the prevailing large-scale weather regimes. The local wind direction at the stations depended on the large-scale conditions but was modified due to complex topography. The NO\textsubscript{x} concentration showed weak correlation for the different stations and depended strongly on the wind direction and atmospheric stability. Conversely, the O\textsubscript{3} concentration was highly correlated among the different measurement sites and was controlled by the long-range atmospheric transport to Svalbard. Lagrangian backward trajectories have been used to examine the origin and path of the air masses during the campaign.
1 Introduction

Fossil fuel combustion and biomass burning create high-temperature conditions, leading to the reaction between atmospheric oxygen and nitrogen present in the fuel and in the air and formation of nitrogen oxides (NO\textsubscript{x} = NO + NO\textsubscript{2}) (Seinfeld and Pandis, 2006). NO\textsubscript{x} emitted locally in the Arctic or transported from midlatitudes may increase the deposition of nitrates (NO\textsubscript{3}\textsuperscript{-}), which act as nutrients and which during climate change may cause changes in the relative abundances of species in nutrient-deficient environments such as lakes in Svalbard (AMAP, 2006). Aerosols, containing particulate nitrate, are formed from the gaseous nitric acid (HNO\textsubscript{3}) produced through the oxidation of nitrogen dioxide (NO\textsubscript{2}) by OH radicals in the presence of sunlight or by the night-time reaction with tropospheric ozone (O\textsubscript{3}) (AMAP, 2006).

High concentrations of NO\textsubscript{x} may lead to regional soil and water acidification and have negative effects on human health (AMAP, 2006). In addition to this, NO\textsubscript{x} are vital for the formation of tropospheric ozone O\textsubscript{3}, which is a harmful air pollutant and a greenhouse gas (IPCC, 2013). The O\textsubscript{3} production and loss depend on the ratios between hydrocarbons/NO\textsubscript{x} and CO/NO\textsubscript{x} (carbon monoxide and nitrogen oxides) and the presence or absence of sunlight. In the absence of sunlight during polar night, O\textsubscript{3} that has been produced within long-range-transported polluted air masses may accumulate in the Arctic. Therefore, the atmospheric lifetime of O\textsubscript{3} may be increased from days in summer to months in winter (Quinn et al., 2008).

Ultraviolet (UV) solar irradiance has a complex influence on NO\textsubscript{x} and O\textsubscript{3} chemistry in the troposphere (Seinfeld and Pandis, 2006). Some of the reactions are only efficient at shorter wavelengths, while other processes occur even at longer wavelengths. The insolation increases rapidly in the Arctic during the springtime transition from polar night to midnight sun, but the UV irradiance is dominated by the UV-A fraction with wavelengths from 315 to 400 nm, while the amount of incoming shortwave UV-B irradiance with wavelengths from about 300 to 315 nm is still minimal in this period (Seinfeld and Pandis, 2006). One of the processes that takes place even under low solar elevation and higher column ozone concentration over the sea-ice and snow-covered surfaces is the photolysis of dihalogens (Simpson et al., 2015). This process is the initial step needed for the heterogeneous photochemical reactions with bromine compounds, promoting the springtime tropospheric O\textsubscript{3} depletion events (Fan and Jacob, 1992; Monks, 2005; Simpson et al., 2015). According to the study performed by Beine et al. (1997a), the background NO\textsubscript{x} values were also lower than normal during such events observed at the Zeppelin station in Svalbard. The reactions with Br species, which result in the oxidation of NO to NO\textsubscript{2} and the removal of NO\textsubscript{2} by the reaction with BrO or OH radical and formation of HNO\textsubscript{3}, were proposed as a possible explanation to this phenomenon. However, lower NO\textsubscript{x} values are also characteristic of the pristine air masses from the remote regions in the high Arctic. In contrast, elevated NO\textsubscript{x} values are observed during the pollution episodes near the local emission sources or when NO\textsubscript{x} are transported to the Arctic from midlatitudes directly or in the form of peroxyacetyl nitrate (PAN), which is further thermally decomposed locally in the Arctic when the air temperature increases in spring (Beine et al., 1997b). Irrespective of the UV irradiance, in the vicinity of large sources of NO, such as cruise ships, the titration of O\textsubscript{3} and formation of NO\textsubscript{2} may be observed (Eckhardt et al., 2013). However, if NO and CO or hydrocarbons are present in sufficient quantities downwind from the emission source and insolation increases (Wallace and Hobbs, 2006), the photolysis of O\textsubscript{3} at wavelengths below 320 nm may lead to production of the OH radical in the presence of water vapour, which may further yield net O\textsubscript{3} production. The UV irradiance also affects the NO/NO\textsubscript{2} ratio. NO\textsubscript{2} dissociates to NO and O in the range of wavelengths from 300 to 370 nm. The photodissociation efficiency reduces gradually at higher wavelengths and vanishes at 420 nm (Seinfeld and Pandis, 2006). A diurnal variation in the background NO/NO\textsubscript{2} ratio has been observed in Svalbard in spring, and the increase in this ratio around noon becomes more pronounced from March to May (Beine et al., 1997b). The efficiency of NO\textsubscript{2} photolysis and formation of NO and O\textsubscript{3} enhances as insolation increases, despite concurrent rapid oxidation of NO by O\textsubscript{3} leading to formation of NO\textsubscript{2}, a second part of the so-called daytime NO\textsubscript{x} null cycle (Wallace and Hobbs, 2006). Thus, both UV-B and UV-A solar irradiance fractions may have influence on the springtime concentrations of NO\textsubscript{x} and O\textsubscript{3} in Svalbard and should be taken into consideration.

Meteorological conditions, such as wind speed and direction, air temperature and humidity, affect the formation of aerosols and efficiency of pollution dispersion and deposition. Synoptic-scale north-easterly wind prevails in the Svalbard region (Adakudlu et al., 2019), but the mesoscale flow is affected locally by topographical channelling and air density gradient from the inland glaciers to the warmer sea. The most pronounced wind direction is along the valleys or fjords towards the coast (Førland et al., 1997): from south-east in Longyearbyen and Ny-Ålesund and from south-south-east in Barentsburg (Fig. 1). Nevertheless, despite the difference in local wind direction in the settlements, there may be common mesoscale meteorological conditions promoting accumulation of locally emitted pollutants in the atmospheric boundary layer (ABL) at all three sites, such as ABL height variation and atmospheric temperature inversion (Dekhtyareva et al., 2018).

The main anthropogenic emission sources on the archipelago are related to electricity and heat production: coal power plants in Barentsburg and Longyearbyen and a diesel-fuelled generator in Ny-Ålesund (Dekhtyareva et al., 2016; Vestreng et al., 2009). The energy demand for heating in Longyearbyen is 2 times higher in winter than in summer due to the lower temperatures in wintertime. On winter days,
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In Svalbard, the snowmobile traffic is allowed only in specific zones created to minimize environmental impact from the usage of motorized vehicles on snow-covered and frozen ground (Klima- og miljødepartementet, 2001). Furthermore, because of complex terrain, most of the snowmobile routes are confined to valleys. Therefore, the pollution dispersion is restricted and strongly affected by local circulation patterns. The amount of pollutants emitted instantaneously by one motorcade may be significant, since tourists and residents usually travel in groups consisting of up to 20 snowmobiles due to safety reasons. Previous studies have shown highly elevated levels of volatile organic compounds along snowmobile tracks (Reimann et al., 2009); however, no measurements of nitrogen oxides have previously been done.

NO\textsubscript{x} concentrations in all three settlements may also be influenced by emissions from ship traffic (Vestreng et al., 2009; Shears et al., 1998), which is the most intensive in summer (Eckhardt et al., 2013; Dekhtyareva et al., 2016), while snowmobiles and power plants are dominant sources of NO\textsubscript{x} in winter and spring seasons.

The main aim of the current article is to combine NO\textsubscript{x} and O\textsubscript{3} data, collected in spring 2017 in Barentsburg, Longyearbyen and Ny-Ålesund, in order to identify specific factors affecting the concentration of measured compounds and define conditions that promote accumulation of local and long-range-transported pollution in all three settlements.

Meteorological in situ and reanalysis data as well as UV, O\textsubscript{3} and NO\textsubscript{x} observations have been used to test the following hypotheses:

- There is a diurnal pattern in concentration of NO\textsubscript{x} at all three stations due to a variable emission rate from the local sources of NO\textsubscript{x}.
- Complex topography determines local circulation, and therefore variation of NO\textsubscript{x} concentration measured at the stations will be dominated by micro- and mesoscale phenomena.
- Local emissions of NO\textsubscript{x} in Ny-Ålesund and Barentsburg affect O\textsubscript{3} concentrations in the settlements.
- Despite the topographically induced features, there are common synoptic meteorological conditions, which have an influence on the concentrations of NO\textsubscript{x} and O\textsubscript{3} in the settlements.

The observational setup at the three stations and methods applied to study various factors of influence on the concentration of measured compounds are introduced in “Materials and methods” (Sect. 2). In the Results (Sect. 3), the NO\textsubscript{x} and O\textsubscript{3} observations from Adventdalen, Barentsburg and Ny-Ålesund are compared, and the influence of large-scale weather regimes on the concentrations of measured compounds at the three stations is identified. In the Discussion (Sect. 4), the results from the 2017 campaign are contrasted to the modelled NO\textsubscript{2}/NO\textsubscript{x} ratio and NO\textsubscript{2} produced through
PAN decomposition and long-term observations from Ny-Ålesund and weather regime and trajectory data are utilized to confirm large-scale circulation and air pollution links. Finally, the Conclusion section summarizes main findings of the paper and implications of the weather regime approach for air pollution studies in Svalbard.

2 Materials and methods

2.1 Measurements in Adventdalen (Longyearbyen)

In the spring season, the main snowmobile route from Longyearbyen to the east coast of Spitzbergen goes along the road through the Adventdalen valley, and therefore there is daily snowmobile traffic nearby the CO₂ laboratory belonging to the University Centre in Svalbard (UNIS CO₂ lab, coordinates: 78.20247° N, 15.82887° E). The station is located at a distance of approximately 5 km to the south-east of the coal power plant and major crossroads in Longyearbyen and thus is representative of monitoring of air pollution from snowmobiles. The chemiluminescence NO/NO₂/NO₃ analyser (model T200) was installed at this laboratory for the period from 23 March to 15 May 2017. The inlet of the sampling hose was secured outside from the window, while the temperature inside the laboratory was kept constant with the help of a thermostat to maintain stable conditions needed for correct functioning of the analyser. The sensor was calibrated weekly using a zero-air generator and a certified NO gas with known concentration (800 ppb), and the hourly average NO₃ data were scaled linearly to eliminate zero drift. The automatic weather station (UNIS AWS) is located nearby the UNIS CO₂ lab, and the data from that station have been used to assess local meteorological conditions.

In addition to the meteorological parameters from the Adventdalen station, data from UV monitors installed at the UNIS roof in Longyearbyen have been used. The sensors SKU 420 UV-A (315–380 nm) and SKU 430 UV-B (280–315 nm), produced by SKYE Instruments, were calibrated on 24 August 2016.

2.2 Measurements in Ny-Ålesund

Continuous NO₃ measurements are performed by the Norwegian Institute for Air Research (NILU) in the framework of the air quality monitoring programme in Ny-Ålesund (Johnsrud et al., 2018). The analyser is installed in the middle of the settlement (coordinates: 78.92470° N, 11.92641° E), 100 m to the north-west of the meteorological station operated by the Norwegian Meteorological Institute and 300 m to the south-south-east of the diesel power plant. Similarly to the measurements in Adventdalen, weekly calibrations with zero air and span gas are performed at the Ny-Ålesund station, and the hourly average NO₃ data are corrected for drift. The hourly O₃ gas monitor data from the Zeppelin observatory, located nearby the mountain top (474 m a.s.l.), 2 km to the southwest of Ny-Ålesund (coordinates: 78.90719° N, 11.88606° E), were used for comparison with the O₃ measurements in Barentsburg. The UV data obtained using the multifilter radiometer GUV 541 at the Sverdrup station in Ny-Ålesund (Gröbner et al., 2010; Schmalwieser et al., 2017) and local meteorological observations from the Zeppelin station were provided by NILU as well. The GUV radiometer is checked and corrected against a travelling reference instrument every year.

In addition to this, daily radiosonde and weekly ozonesonde data from the French–German AWIPEV research station in Ny-Ålesund have been used. Since temperature inversion may be an important factor promoting accumulation of local pollution in the atmospheric boundary layer, the method for its detection in the radiosonde vertical profiles, described by Dekhtyareva et al. (2018), has been applied: the days when the temperature was increasing with height by more than 0.3 °C in the lowest 500 m were defined as days with temperature inversions. In order to compare the O₃ sonde measurements with ground-level observations, the O₃ mixing ratio in units of parts per billion per volume (ppbv) were calculated from the O₃ partial pressure and atmospheric pressure measured by the radiosonde (Seinfeld and Pandis, 2006).

Daily radiosonde launches were operated at the AWIPEV station (AWI), using Vaisala RS92 radiosondes until April 2017 (Maturilli and Kays, 2017) and Vaisala RS41 radiosondes afterwards. In this study, we apply radiosonde data for March 2017, post-processed according to the principles of the GCOS Reference Upper-Air Network (GRUAN; Immler et al., 2010). The RS41 data for April–May 2017 are processed with the manufacturer’s software.

2.3 Measurements in Barentsburg

The Russian Arctic and Antarctic Research Institute (AARI) performs the measurements in Barentsburg independently in the frame of the air quality monitoring programme. The equipment installed in the settlement includes chemiluminescence NO₃ AC32M and UV photometric O₃ O₃/42M analysers produced by Environment S.A. and the Vaisala HydroMet system MAWS201. The observational site is located on the narrow terrace 40 m above sea level (coordinates: 78.06070° N, 14.21718° E) and 600 m to the north-east of the coal power plant. The analysers continuously gather the data and transmit them to the laboratory facility of the Russian Scientific Centre in Barentsburg. The data with 20 min time resolution were averaged to obtain hourly data. The analysers were installed and initially calibrated by the manufacturer’s accredited specialists in December 2016. After the installation, zero control was performed regularly using the NO₃ analyser’s built-in automatic zero-air function for the correction of zero drift lines. The maintenance of the converter filter was done at the frequency recommended by the manufacturer. However, in contrast to the equipment in Ny-Ålesund
and Longyearbyen, the NO$_x$ and O$_3$ analysers in Barentsburg were not calibrated manually during the field campaign. Therefore, the data from this station may be prone to drift. This is especially important to keep in mind when studying NO$_x$ concentrations, since the NO$_x$ values are usually very low in the remote Arctic environment (Dekhtyareva et al., 2016). On the other hand, the UV O$_3$ monitor is more stable and does not demand as frequent calibration as chemiluminescence instruments (Williams et al., 2006), and thus data from this instrument are more reliable.

2.4 Methods to study the effect of meteorological conditions on the concentration of measured compounds

Previous studies showed the influence of large-scale weather phenomena on long-range-transported and local air pollution. Eckhardt et al. (2003) demonstrated how positive and negative phases of the North Atlantic Oscillation control long-range transport of air pollution to the Arctic. The modelling study of Ménégoz et al. (2010) introduced four weather regimes based on ERA40 data and described the aerosol budget variations associated with different regimes and feedback of aerosol distribution on the weather regime persistence. Ibrahim et al. (2021) classified nine weather regimes using self-organizing maps and cluster analysis of principal components and investigated the influence of the prevailing large-scale meteorological conditions on the air quality in Berlin. In the current work, we apply Christian Grams’ weather regime classification that is based on the 6-hourly ERA-Interim data. This classification was previously used to investigate the frequency of poleward moisture transport events by atmospheric rivers (Pasquier et al., 2019) and southward transport of Arctic air during cold-air outbreaks (Papritz and Grams, 2018) and for assessment of Europe’s wind power output (Grams et al., 2017). Thus, this approach is suitable for the study of long-range transport of air masses and local dispersion efficiency that depends on atmospheric stability and wind speed. The following three-step procedure has been implemented to assess the effect of the prevailing synoptic meteorological situation on long-range transport of pollutants and on the local meteorological conditions affecting concentrations of O$_3$ and NO$_x$ in Svalbard.

Firstly, the whole measurement period was divided into sub-periods based on the prevailing large-scale atmospheric circulation patterns. In the climatological mean, the large-scale conditions are characterized by weak ridging of absolute geopotential height at 500 hPa over the eastern North Atlantic and westerly upper level flow over Svalbard. Such a regime is placed in the “no regime” category in the Grams et al. (2017) classification (their Fig. S1h). The deviations from these mean conditions are classified into seven distinct weather regimes that represent the variation of the large-scale circulation patterns over the North Atlantic and European region.

Secondly, the hourly meteorological data, O$_3$ concentration and planetary boundary layer height (PBL) from the global ERA5 reanalysis data set with 31 km spatial resolution (Hersbach et al., 2020) were used to investigate the prevailing large-scale weather conditions for the identified sub-periods. The ERA5 O$_3$ mass mixing ratio is estimated based on the assimilated satellite observations and external 2-D photochemical model (Park et al., 2020). The PBL in the ERA5 data set is defined by the height when the Richardson number for the adjacent vertical model layers exceeds a critical value of 0.25, and the air becomes stably stratified (European Centre for Medium-Range Weather Forecasts, 2017).

Thirdly, the FLEXible PARTicle (FLEXPART) V8.2 air parcel trajectory data set was utilized for the same sub-periods to study the long-range atmospheric transport to Svalbard. Previously, FLEXPART data have been used to investigate long-range transport of black carbon and sulfates (Hirdman et al., 2010a, b) and mercury and O$_3$ (Hirdman et al., 2009) in the Arctic. The current FLEXPART data set contains a 3-dimensional Lagrangian dispersion simulation (Stohl et al., 2005) with 5 million air parcels (Läderach and Sodemann, 2016; Fremme and Sodemann, 2019) driven with the meteorological data from the ERA-Interim reanalysis with a spatial resolution of approximately 80 km and temporal resolution of 6 h (Dee et al., 2011). The 10 d backward trajectories starting within the lowest 500 m above the ground in the region covering Ny-Ålesund, Longyearbyen and Barentsburg (from 77.5 to 79.5° N and from 10 to 20° E) were extracted from this data set. The trajectory model data were combined with BrO total column data derived from GOME-2 (ir)radiance satellite observations (AC SAF, 2017) to identify regions with elevated concentration of this O$_3$-depleting substance.

Additionally, to study long-range transport of extremely O$_3$-depleted or O$_3$-enriched air masses, the following procedure has been implemented to detect O$_3$ decrease and increase events occurring simultaneously in Barentsburg and at the Zeppelin station:

1. Since the distance between the Zeppelin observatory and Barentsburg is more than 100 km, a time lag in correlation between the data from the two stations is expected. The acceptable time lag has been defined based on the lagged linear correlation between the data sets. The maximum time lag for which the correlation coefficient is higher than or equal to the coefficient calculated for the zero-hour lag is defined as the maximum allowable time lag.

2. Applying the extreme definition of air quality stated in Porter et al. (2015), O$_3$ levels below the 5th quantile and above the 95th quantile were found separately for the Barentsburg and Zeppelin measurements to define severe depletion and increase events, respectively.
3. Continuous episodes were defined for the periods where the time difference between consecutive event points is less than 3 h.

4. Minimum (maximum) O$_3$ concentrations within each event were defined. The time of minimum (maximum) concentration was chosen for both stations, and if the difference between them was less than the acceptable time lag, the events at both stations were classified as one joint event.

The backward air mass ensemble trajectories have been simulated using the Hybrid Single Particle Lagrangian Integrated Trajectory (HYSPLIT) model for these joint events for 240 h back in time to identify the source regions of the air masses (Stein et al., 2015). This 10 d simulation time has been chosen as a compromise between the average lifetime of tropospheric O$_3$, which may be 3 to 4 weeks (Christiansen et al., 2017), and the uncertainty of modelled air mass trajectories that increases with travelled distance (Freud et al., 2017). The standard ensemble simulation with 27 members was calculated in the READY system by offsetting the Global Data Assimilation System (GDAS) meteorological data with a 0.5° horizontal and vertical dimensions to take into account the uncertainty of the trajectory forecast (Rolph et al., 2017).

2.5 Methods to study the effect of local pollution in Ny-Ålesund and in Barentsburg on the measured O$_3$ concentrations

Previously, the only collocated NO$_x$ and O$_3$ measurements from Ny-Ålesund were performed at the Zeppelin station from February to May 1994. The results were published in Beine et al. (1996). In that study, the combination of NO$_x$ data and concentration of particles with diameter below 10 nm, atmospheric stability and wind direction was used to identify possible local pollution events. In spring 1994, the local pollution was detected at the Zeppelin station during 6.4 % of the measurement time, and the number of these events increased with increased insolation in May. As there were no O$_3$ data from Ny-Ålesund available, the O$_3$, CO and differential mobility particle sizer (DMPS) data from the Zeppelin station were used to study the influence of local NO$_x$ emissions in Ny-Ålesund on the O$_3$ concentration. CO indicates the presence of other pollutants emitted simultaneously in the process of fossil fuel burning, and although the correlation between NO$_x$ and CO concentration in the plumes depends on the engine and fuel type, age of the plume and environmental conditions (Li et al., 2015), we expect elevated CO concentrations in the fresh plumes arriving at the Zeppelin station. Therefore, a local pollution effect was defined for O$_3$ measurements at the Zeppelin station when all four conditions were fulfilled:

1. The wind direction measured both in Ny-Ålesund and at the Zeppelin station was northerly (above 270° or below 90°) since the diesel power plant is located 300 m to the north–north-west of the NO$_x$ monitor in Ny-Ålesund and 2 km to the north–north-east of the Zeppelin station.

2. NO$_x$ concentrations were above the mean value in Ny-Ålesund.

3. The concentration of particles with diameter below 10 nm had a strong increase (above 95 percentile value for the whole campaign).

4. CO concentrations observed at the Zeppelin station were above the mean value, indicating the possible impact of local pollution.

To assess how the NO$_x$ emissions in Barentsburg affect the local O$_3$ concentration there, the NO$_x$ and O$_3$ data from the Barentsburg station have been compared. Positive anomalies in O$_3$ concentration were found for the same wind directions where increased NO$_x$ concentrations were observed, but this may be due to higher concentrations of O$_3$ in air masses, which were transported to Svalbard from the south and south-west. Since there are multiple sources of local pollution in Barentsburg (coal power plant, ships and cars), another method has been implemented:

1. The hours when NO$_x$ concentrations were above average in Barentsburg have been defined.

2. O$_3$ values for these hours in the original and in the 6-hourly smoothed data series from Barentsburg have been compared.

3 Results

3.1 Comparison of NO$_x$ and O$_3$ observations from Adventdalen, Barentsburg and Ny-Ålesund

The hourly NO$_x$ and O$_3$ concentrations and meteorological data from all three stations are shown in Figs. 2 and 3. Nine distinct large-scale weather regimes have been identified for the campaign period (marked with numbers from I to IX). A detailed description of the weather regimes is given in Sect. 3.2 of the current paper.

Despite alteration in the large-scale circulation, little variability was found in CO data from the Zeppelin station (solid olive line in the Fig. 2c) and O$_3$ values in ERA5 reanalysis (orange line in the Fig. 2c). The threshold of CO median value ± median absolute deviation (132.6±6.7 ppb) is shown by the dashed olive line in the Fig. 2c. There were no sharp peaks in the concentration of this gas, indicating that there was little influence of local pollution on the measurements at the mountain station, but the levels of this compound were stably high in the beginning of the campaign and showed a gradual decline from the end of March to the middle of May. This is a response to increased insolation throughout the fieldwork period (from 0.03 to 0.3 W m$^{-2}$ and from 7.9
Figure 2. (a) The hourly average air temperature and relative humidity measured in Ny-Ålesund (NYA) and at the Zeppelin station (Zep). (b) Atmospheric pressure and wind speed measured in Ny-Ålesund and at the Zeppelin station. (c) NO\textsubscript{x}, CO and O\textsubscript{3} concentrations in Ny-Ålesund, at the Zeppelin station and in the ERA5 reanalysis for the nearest grid point the coordinates of Ny-Ålesund for the period from 23 March to 16 May 2017. The dashed black lines represent time frames of the different weather regimes. The dashed blue line shows 5th and 95th quantiles of the O\textsubscript{3} concentration.

Figure 3. (a) The hourly average air temperature and relative humidity measured in Adventdalen (Adv) and Barentsburg (BBG). (b) Atmospheric pressure and wind speed measured in Adventdalen and Barentsburg. (c) NO\textsubscript{x} and O\textsubscript{3} concentrations in Adventdalen, in Barentsburg and in the ERA5 reanalysis for the nearest grid point to the coordinates of Barentsburg for the period from 23 March to 16 May 2017. The dashed black lines represent time frames of the different weather regimes. The dashed light blue line shows 5th and 95th quantiles of the O\textsubscript{3} concentration.
to 28.9 W m\(^{-2}\) UV-B and UV-A irradiation observed in Ny-Ålesund, respectively), as CO is rapidly oxidized by the OH radical produced in the O\(_3\) photolysis. A pronounced decline in CO concentration and sharp drop in O\(_3\) values in ERA5 reanalysis may be observed in sub-period VIII. The O\(_3\) concentration in the ERA5 reanalysis exceeds the values observed at the Zeppelin station (dark blue line in the Fig. 2c) most of the time except this sub-period. Thus, the O\(_3\) ERA5 reanalysis data are not representative of the regional Arctic processes or short-term variability in long-range transport but show strong sensitivity to photochemical destruction.

There is a weak statistically significant positive correlation between NO\(_x\), NO\(_2\) and NO\(_x\) values measured in Adventdalen and in Ny-Ålesund (the Pearson correlation coefficients are \(r_{\text{NO}} = 0.13, r_{\text{NO}_2} = 0.15\) and \(r_{\text{NO}_x} = 0.13; p < 0.0001\) for all compounds). Conversely, no correlation is present with NO\(_x\) data from Barentsburg. Low correlation between the NO\(_x\) values at the three stations indicates the importance of local emission sources and micrometeorology (wind channelling and spatial variation in atmospheric stability) rather than synoptic meteorological conditions. The background NO\(_x\) concentrations observed in Svalbard in previous studies (Beine et al., 1997a, b) using different measurement techniques are below 0.4 ppb, and thus, the natural variability in NO\(_x\) values due to long-range transport to Svalbard would be undetected in the NO\(_x\) data sets presented in the current study.

The Barentsburg O\(_3\) data contain some abrupt peaks with magnitude of up to 9 ppbv and duration of just 1 h (light blue line in Fig. 3c), while they are absent in the Zeppelin O\(_3\) data. Indeed, the Barentsburg station is located inside the settlement, and the O\(_3\) data there are more prone to be influenced by local NO\(_x\) pollution, while the Zeppelin station is situated far from the local emission sources. In order to investigate the significance of this local impact, a 6 h moving average filter has been applied to the O\(_3\) Barentsburg data, and the results are shown with a dark blue line in Fig. (3c). The smoothed and original O\(_3\) data from Barentsburg have been compared statistically: both the two-sided Wilcoxon rank sum (WRS) test and the \(t\) test show that the application of the low-pass filter on the O\(_3\) data from Barentsburg does not result in significant change in the concentration distribution. The correlation between O\(_3\) concentrations at the Zeppelin station and in Barentsburg is moderate (Pearson correlation coefficient \(r = 0.69\) both for smoothed and unsmoothed data; \(p < 0.0001\)). This indicates that O\(_3\) concentrations at both stations are highly influenced by the meteorological conditions on the synoptic scale, and local impacts are of minor importance.

We have applied methods described in Sect. 2.5 to define the effect of local NO\(_x\) emissions in Ny-Ålesund and Barentsburg on the O\(_3\) concentrations in the settlements. As a result, 5% of the O\(_3\) data from the Zeppelin station might have been influenced by the local pollution from Ny-Ålesund, and the statistically significant \((p < 0.0001)\) decrease in the O\(_3\) mean (31.6 vs. 36.1 ppb) and median (34.4 vs. 38.0 ppb) concentrations has been revealed for this group. However, northerly wind that may transport local pollution from Ny-Ålesund also brings air masses which have a lower O\(_3\) background value. Therefore, when one compares potentially locally polluted air masses with the background air masses coming from the north, the difference in mean and median O\(_3\) concentrations becomes statistically insignificant, 31.6 vs. 33.3 ppb and 35.6 vs. 34.4 ppb, respectively. Following the method of Beine et al. (1996) for the local pollution event detection, the concentration of particles with a diameter of 10 nm routinely measured by the DMPS at the Zeppelin station and a threshold of > 95 percentile have been used to identify peaks in concentration of newly formed particles. Similarly to the results of Beine et al. (1996), the peak events were detected at the Zeppelin station only in the second part of the 2017 measurement campaign (from 24 April to 13 May). The northerly wind direction was present only during 12 out of 45 h with peak particle concentration; however, none of these cases was characterized by increased CO concentration at the Zeppelin station. Thus, these peaks in concentration of small particles might have been related to natural rather than anthropogenic emission sources. Indeed, Heinzenberg et al. (2017) described the offset in new particle formation towards late spring and summer when biological emissions become important sources for this process. Therefore, both statistical comparison of the O\(_3\) concentrations in clean and potentially polluted air masses mentioned above and the absence of coinciding peaks in particle concentration and CO concentration indicate that the O\(_3\) observations at the Zeppelin station were not significantly affected by local NO\(_x\) pollution during the 2017 campaign.

The difference between the original and smoothed O\(_3\) data from Barentsburg varies from −19% to 11% of the smoothed value, and there is a moderate negative correlation between the magnitude of NO\(_2\) peak and the reduction in O\(_3\) concentration \((r = −0.65, p < 0.0001)\). Despite this sensitivity of O\(_3\) concentration to local pollution in Barentsburg, the median NO\(_x\) concentrations observed there were low, and average reduction of O\(_3\) concentration in comparison to the smoothed values was only 1%. This effect is not statistically significant, and therefore other factors, such as variation in concentrations within long-range-transported air masses, may be more important for explanation of difference between the O\(_3\) Zeppelin and Barentsburg data sets.

The comparison of the vertical O\(_3\) data from the O\(_3\) sondes (dark blue squares in Figs. 2c and 3c) from Ny-Ålesund and the ground-based measurements at the Zeppelin station and in Barentsburg reveals that the discrepancy in the data between the two stations may be explained by the fact that the stations are located at different heights and measure air masses with an uneven distribution of O\(_3\) in the lowest 500 m. If contrasting the closest point to the sounding time in the observations made in Barentsburg and in Ny-Ålesund, similar tendencies as in the O\(_3\) sonde data may be observed. For ex-
ample, there is a significant difference between the data from Barentsburg and Ny-Ålesund (33.74 ppb vs. 38.88 ppb) for the measurement closest to the time of sounding on 10 May, and an increase of O\textsubscript{3} concentration with height between 1000 hPa (50 m) and 950 hPa (500 m) is noticeable in the sounding data as well (Fig. 4a). One can see in the potential temperature profiles (Fig. 4b) that the pronounced atmospheric inversion tends to be noticeable in the O\textsubscript{3} sondes profiles as well. For example, a simultaneous increase in O\textsubscript{3} concentrations and potential temperature is pronounced at the level of 850 hPa (1300 m) on 26 April, 860 hPa (at 1200 m) on 5 April and 900 hPa (at 1000 m) on 12 April (Fig. 4a and b). The O\textsubscript{3} concentration in the ERA5 profiles was overestimated and showed little variability in the lowermost layer, except the profile for 10 May (dashed dark grey line in the Fig. 4a) when the reanalysis and observational values coincided for 1000 hPa level, but the O\textsubscript{3} concentration in ERA5 was underestimated by almost 40% at the height of 925 hPa (Fig. 4a). The virtual potential temperature profiles in reanalysis resemble O\textsubscript{3} sonde profiles closely (Fig. 4b). The PBL height from ERA5 reanalysis (marked with stars in the Fig. 4b) was above the low-level temperature inversions detected on 5 and 12 April and below the level of the most pronounced virtual temperature inversion in the O\textsubscript{3} sondes profiles.

In addition to hourly values, average concentrations of measured compounds have been calculated for each hour of the day. The diurnal variation in NO, NO\textsubscript{2} and O\textsubscript{3} concentrations and potential temperature is pronounced at the level of 850 hPa (1300 m) on 26 April, 860 hPa (at 1200 m) on 5 April and 900 hPa (at 1000 m) on 12 April (Fig. 4a and b). The O\textsubscript{3} concentration in the ERA5 profiles was overestimated and showed little variability in the lowermost layer, except the profile for 10 May (dashed dark grey line in the Fig. 4a) when the reanalysis and observational values coincided for 1000 hPa level, but the O\textsubscript{3} concentration in ERA5 was underestimated by almost 40% at the height of 925 hPa (Fig. 4a). The virtual potential temperature profiles in reanalysis resemble O\textsubscript{3} sonde profiles closely (Fig. 4b). The PBL height from ERA5 reanalysis (marked with stars in the Fig. 4b) was above the low-level temperature inversions detected on 5 and 12 April and below the level of the most pronounced virtual temperature inversion in the O\textsubscript{3} sondes profiles.

The NO\textsubscript{2}/(NO + NO\textsubscript{2}) ratio is quite high in Adventdalen and in Barentsburg and exhibits diurnal variation, while it is much lower in Ny-Ålesund, and there is no statistically significant difference between its day and night values. This may be explained by the fact that the measurement station in Ny-Ålesund was located much closer to the diesel power plant, a constant source of fresh NO\textsubscript{2} emissions, where the NO\textsubscript{2}/O\textsubscript{3} ratio is much lower irrespective of the time of the day (Beine et al., 1996). The lowest hourly NO\textsubscript{2}/O\textsubscript{3} ratio of 0.29 and the highest peak of NO\textsubscript{2} were observed in Ny-Ålesund at 17:00 UTC 28 April (Fig. 2). The concentration of NO and NO\textsubscript{2} was 87.8 and 16.4 ppb, respectively, which indicates the presence of a strong emission source, for example snowmobiles, in the immediate vicinity from the station. Since this was a single NO\textsubscript{2} peak in the data, the NO\textsubscript{2}/NO\textsubscript{3} ratio was unusually low, and the meteorological conditions were untypical for pollution accumulation in Ny-Ålesund (south-easterly wind with moderate speed of 5.3 m s\textsuperscript{-1}); hence this value has been excluded from further statistical analysis.

NO is a primary product of fossil fuel combustion (Seinfeld and Pandis, 2006; Arya, 1999), and a higher NO/NO\textsubscript{3} ratio is expected close to the emission source. The station in Adventdalen is located at a distance of 5 km from the coal power plant, and snowmobile traffic there is a temporarily emission source present mostly during daytime. In contrast, measurement stations in Barentsburg and Ny-Ålesund are located near the power plants, constantly releasing combustion products at a variable rate. Thus, it is noticeable in Adventdalen that the NO concentration is close to zero during the night (dark blue bar in Fig. 5a) in the absence of fresh traffic emissions and photochemical conversion of NO\textsubscript{2} to NO. As the traffic intensity increases during the day, the NO concentration rises, however, and so does the NO\textsubscript{2} concentration (red bar in Fig. 5a), since there is a rapid conversion of NO to NO\textsubscript{2} by the reaction with O\textsubscript{3}.

There is a slight increase in the O\textsubscript{3} values in Barentsburg (grey line in Fig. 5b) during daytime. In contrast, a slight decrease in daytime O\textsubscript{3} concentration is observed at the Zeppelin station. The discrepancy in the diurnal O\textsubscript{3} profiles from the two stations may be explained by the difference of the altitude of each of the two stations and response of the measurements to the boundary layer dynamics. The ERA5 data show diurnal variation in the PBL height for both locations with highest average values of 462 and 293 m, in Barentsburg and Zeppelin, respectively, for 13:00 UTC. The Barentsburg station is located at the altitude of 50 m a.s.l. and samples air within the ABL. During the daytime, the vertical mixing between the atmospheric surface layer and the air masses aloft enhances, and the boundary layer height increases. This mixing process may enrich the surface layer with O\textsubscript{3}. A similar influence of convection on replenishing the O\textsubscript{3} in the Arctic ABL after the depletion events has been described in the work of Moore et al. (2014). In contrast, the Zeppelin station is located at the altitude of 474 m a.s.l. and mostly samples air from the free troposphere with higher O\textsubscript{3} concentration, and thus the data from this station do not exhibit similar diurnal variation as the Barentsburg station. However, the magnitude of these effects is small, and according to the t test and the WRS test, there is no statistically significant difference between the night-time and daytime O\textsubscript{3} concentrations measured at the stations (Table 1).

The average NO and NO\textsubscript{2} concentrations measured at the stations are distributed unevenly over the wind directions. In Adventdalen, the average wind speed was 5.1 ± 3.1 m s\textsuperscript{-1}. The south-easterly wind dominated during the field campaign (Table 2), and there was no significant difference between the daytime and night-time observations. The highest average daytime NO and NO\textsubscript{2} concentrations were observed when the wind was from north-east and south-east in Adventdalen (Fig. 6). To test if the size of snowmobile motorcade has an influence on the NO\textsubscript{2} concentration in Adventdalen, manual observations of the number of snowmobiles were done in 19 d. In general, the effect of a large number of snowmobiles was only noticeable in the NO\textsubscript{3} data when there was low wind speed. For example, in the evening of 1 May 2017, the wind speed was 1.9 m s\textsuperscript{-1}, and the NO\textsubscript{2}
concentration increased sharply to 7.3 ppb due to 21 snowmobiles passing by the station. A group of a similar size was passing by in the evening of 2 May 2017, but the effect on NO\(_2\) values was 3 times lower as the wind speed was higher (4.0 m s\(^{-1}\)). The maximum hourly NO\(_2\) concentration of 11.4 ppb was measured during the Easter holiday on 13 April 2017. In that day, the combination of increased recreational traffic and mild weather conditions (wind speed below 1 m s\(^{-1}\) and air temperature \(-8^\circ\)C) led to accumulation of concentration 13 times higher than daytime hourly average measured during the field campaign. Such low wind speed is untypical for the wind regime in Adventdalen, where normally ventilation is sufficient to effectively disperse NO\(_x\) emitted by the usual amount of motorized traffic. The highest
Table 1. Measurement results from Adventdalen, Barentsburg and Ny-Ålesund. The two-sided $t$ test compares daytime and night-time concentrations at each station and checks if there is a significant difference in mean values for these two groups. The two-sided WRS test compares daytime and night-time concentrations at each station and checks if there is a significant difference in median values for these two groups. Pairs with significant ($p < 0.05$) $t$ and WRS test results are shown with bold font.

| Compound and station | Daytime mean value | Night-time mean value | $p$ value, $t$ test | Daytime median value | Night-time median value | $p$ value, WRS test |
|----------------------|-------------------|----------------------|---------------------|----------------------|------------------------|---------------------|
| NO (ppb):            |                   |                      |                     |                      |                        |                     |
| Adventdalen          | 0.30              | 0.00                 | 0.000               | 0.12                 | 0.00                   | 0.000               |
| Barentsburg          | 0.15              | 0.08                 | 0.000               | 0.03                 | 0.01                   | 0.000               |
| Ny-Ålesund           | 1.29              | 0.52                 | 0.001               | 0.14                 | 0.02                   | 0.000               |
| NO$_2$ (ppb):        |                   |                      |                     |                      |                        |                     |
| Adventdalen          | 0.94              | 0.41                 | 0.000               | 0.49                 | 0.28                   | 0.000               |
| Barentsburg          | 0.39              | 0.54                 | 0.068               | 0.00                 | 0.00                   | 0.099               |
| Ny-Ålesund           | 0.82              | 0.33                 | 0.000               | 0.15                 | 0.03                   | 0.000               |
| NO$_2$/ (NO + NO$_2$) ratio: | | | | | | |
| Adventdalen          | 0.80              | 0.83                 | 0.009               | 0.82                 | 0.85                   | 0.000               |
| Barentsburg          | 0.72              | 0.80                 | 0.000               | 0.78                 | 0.89                   | 0.000               |
| Ny-Ålesund           | 0.61              | 0.63                 | 0.369               | 0.64                 | 0.63                   | 0.335               |
| O$_3$ (ppb):         |                   |                      |                     |                      |                        |                     |
| Barentsburg          | 35.64             | 34.81                | 0.139               | 37.33                | 35.74                  | 0.051               |
| Zeppelin             | 35.55             | 36.14                | 0.203               | 37.18                | 38.28                  | 0.057               |

Average night-time NO$_2$ concentrations were detected when the wind was from north-west, which reveals possible influence of the coal power plant. The average night-time concentrations of NO were very low, regardless of the wind direction.

Figure 7a and b illustrate distribution of NO and NO$_2$ concentrations over wind directions in Ny-Ålesund and Barentsburg, respectively. South-easterly wind with average speed of 3.6 and 3.9 m s$^{-1}$ in daytime and night-time, respectively, was dominating in Ny-Ålesund. However, the highest average NO$_x$ concentrations in Ny-Ålesund were measured when the wind was coming from the north (Fig. 7a). This points clearly to the local diesel power plant being the main emission source. Similar results regarding the influence of the local power plant in Ny-Ålesund on NO$_x$ concentrations were presented in Dekhtyareva et al. (2016) and Johnsrud et al. (2018). During the field campaign, the prevailing wind in Barentsburg was from south and south-east, with an average speed of 2.5 m s$^{-1}$ and from south-east and east with mean speed of 2.3 m s$^{-1}$ in daytime and night-time, respectively. The NO$_x$ concentrations measured there were much lower and more evenly distributed over different wind directions than in Ny-Ålesund (Fig. 7b). The coal power plant operates day and night and, in the light wind conditions, may contribute to accumulation of local pollution in the settlement even in the absence of south-westerly wind.
3.2 Influence of large-scale weather regimes on the concentrations of measured compounds at the three stations

An overview of median concentrations of measured compounds and prevailing local meteorological conditions for the nine sub-periods defined based on the prevailing weather regimes is given in Table 2. The wind directions observed at the stations during each of the sub-periods have been sorted into 16 bins with a 22.5° interval. The three main wind directions for each sub-period and for the whole campaign are stated with letters in Table 2. A detailed analysis of the meteorological conditions prevailing during each of the weather regimes and their influence on the concentration of the compounds measured at the three station is given below.

The “no regime” conditions were present for almost 20 d or 37 % of total campaign duration. The sub-period I lasted for 14 d and was characterized by the lowest median temperatures and UV irradiance at all stations and by synoptic-scale north-easterly flow (Fig. 8a). However, the conditions were inhomogeneous for this period: there were two quick passing cyclones on the 26 March and 3 April (pressure drops in Figs. 2b and 3b) that led to an increase in local air temperatures and wind speeds. In the sub-period I, the temperature inversions were observed in 46 % of the radiosonde profiles from Ny-Ålesund, but the median inversion strength was below 0.95 °C (median for the whole campaign).

The sub-periods II–V were characterized by Atlantic ridge (AR) and Scandinavian trough (ScTr). These regimes are characterized by the varying degree of geopotential height ridging over the North Atlantic at 500 hPa. During the AR regime, an upper-level north-westerly flow prevails (Grams et al., 2017). The strongest wind speed was observed in Adventdalen during the AR regimes, since the synoptic-scale lower-level flow (Fig. 8b and d) was parallel to the Adventdalen valley (Fig. 6). During the transition from AR to ScTr, the upper-level ridge weakened and shifted southwards (Grams et al., 2017), as did the cyclonic systems on the lower level (Fig. 8c and e). The AR and ScTr regimes were characterized by the PBL height being below the median value for the campaign at all sites. The temperature inversions were observed for the regimes II, III and IV with an inversion frequency of 67 %, 57 % and 13 %, respectively, and the inversion strength was above the median for the campaign.

The sub-period VI was a 3 d “no regime” transition between the AR and ScTr and two blocking regimes: Scandinavian blocking (ScBL) and Greenland blocking (GL). This sub-period was characterized by the synoptic-scale westerly wind bringing warm Atlantic air over Svalbard (Fig. 8f), increasing local temperature and PBL height and adding the westerly component to the wind direction at all stations.

During the sub-period VII (ScBL), the positive geopotential height anomaly was located over northern Scandinavia, and part of the upper-level flow was deflected poleward around the blocking anticyclone (Grams et al., 2017). The anticyclonic movement was pronounced in the lower-level flow (Fig. 8g), and the synoptic-scale north-westerly flow prevailed over the western part of Svalbard. The local wind speed, temperature and PBL height decreased.

The sub-period VIII (GL) was characterized by the strong positive anomaly in the geopotential height at 500 hPa over Greenland and the prevailing upper level north-westerly wind (Grams et al., 2017). The lower-level blocking over Greenland promoted north-easterly flow over Svalbard (Fig. 8h).
The sub-period IX ("no regime") was characterized by the strong anticyclone over the Barents Sea, that led to pronounced transport of warm Atlantic air with southerly flow to Svalbard (Fig. 8h). No temperature inversions were detected in the radiosonde data from Ny-Ålesund for the sub-periods VII, VIII and IX.

The elevated NO\textsubscript{x} concentrations were observed in Ny-Ålesund and Adventdalen during the sub-periods I–V. This may be explained by the enhanced accumulation of locally emitted NO\textsubscript{x} in the ABL due to suppressed vertical mixing on cold days associated with the AR and ScTr regimes. In Adventdalen, NO\textsubscript{x} concentrations are not dependent on the wind direction (Fig. 6 and Table 2), and east-south-easterly and south-easterly are the dominant wind directions for all regimes except for the period VI. The highest median values of NO\textsubscript{x} were observed during the periods with the lowest PBL height, ScTr regimes. During the periods VI and VII, the PBL height increased, but westerly and west-north-westerly wind might have brought pollution from the coal power plant and the town of Longyearbyen to the Adventdalen valley (Fig. 6). In Ny-Ålesund, the highest median NO\textsubscript{x} value was observed in sub-period I due to the presence of north-north-westerly wind that brought the plume from the power plant to the measurement station (Fig. 7a).

In contrast to Adventdalen, the boundary layer height and cold temperatures played a secondary role for the NO\textsubscript{x} concentrations in Barentsburg, and the controlling factor was south-westerly component of the wind for the most polluted periods. The highest median NO\textsubscript{x} values were detected during the sub-period VI, when south-south-westerly wind was dominating. The major emission sources in Barentsburg are located on the seashore, and warmer marine air from west and south-west may bring local pollution to the station situated on the hill above these sources (Fig. 7b). However, the second highest median NO\textsubscript{x} value was observed for the sub-period II, the period with the lowest PBL height and easterly wind at this station. The wind direction was not from the coal power plant, but the wind speed was very low, and thus, the local pollution could accumulate in the ABL if a strong inversion was present aloft.

The O\textsubscript{3} data show similar variability for the Zeppelin station and Barentsburg. The concentrations below the campaign’s median were observed for the sub-periods V, VII and VIII. The FLEXPART 10\textsuperscript{d} backward trajectory probability contours show that for these sub-periods, the air masses passed over the region north of Svalbard where the concentration of BrO was elevated (Fig. 9e, g, h). Conversely, the sub-periods III, IV, VI and IX, with O\textsubscript{3} concentration above the median at both stations, are characterized by the air masses arriving from the south-east, east, west and south-west (Fig. 9c, d, i), respectively. In the sub-period I, 24% of the data from Barentsburg were missing. This sub-period’s concentrations at the Zeppelin station were slightly higher than the campaign’s median, despite the most significant O\textsubscript{3} depletion episode occurring on 31 March–1 April (Fig. 2).

The trajectory contours show possible influence of the local depletion in the Svalbard region (Fig. 9a). In the sub-period II, 67% of the data from the Zeppelin station were missing (Fig. 2). The O\textsubscript{3} concentration in Barentsburg was above the median for this sub-period, and the trajectory data show the air masses arriving from the south-east (Fig. 9b). As in previous studies of Hirdman et al. (2009), the downward transport of O\textsubscript{3}-enriched air masses from higher altitudes played a significant role during the 2017 campaign. The percentage of trajectory points reaching elevations above 2000 m was highest for the sub-periods III, VI and IX (27%, 33% and 24% of the total number trajectory points for each sub-period respectively). In contrast, during the sub-period VIII with the lowest O\textsubscript{3} concentration at both stations, the percentage of elevated trajectory points was minimal, only 4%. One can also see that the percentage of elevated trajectories varies for the same type of weather regime and determines importance of the downward air mass transport for the measured surface O\textsubscript{3} concentrations in different sub-periods (e.g. ScTr regime in Fig. 9c and e and Table 2).

Two joint extreme O\textsubscript{3} depletion events (31 March and 6 May 2017) and three increase events (13 April, 28 April and 3 May 2017) have been detected (Figs. 2 and 3). The HYSPLIT trajectory analysis shows that these O\textsubscript{3} depletion events occurred when the cold air masses from the central Arctic reached Svalbard. The trajectory for the strongest depletion episode is shown in Fig. (10a). The concentration of O\textsubscript{3} in the Arctic air masses may be lower because of a lack of sunlight and O\textsubscript{3} precursors such as NO\textsubscript{x}, hydrocarbons and CO needed for the O\textsubscript{3} formation. Further depletion may have occurred due to photochemical reactions with bromine species over the sea ice in the period from 30 March 2017 at 10:00 to 31 March 2017 at 17:00 when the trajectories passed the region with elevated BrO concentration between 80°N and 85°N. The simulated median sun flux was quite low (67 W m\textsuperscript{-2}) but probably sufficient enough to support the halogen-induced O\textsubscript{3} destruction, which might occur even under low-light conditions (Simpson et al., 2015). The trajectories for the increase events revealed southerly origin of the air masses, but source regions were different for all three cases. In the first case, air masses were arriving from the northern part of Russia and in the second one from North America and Iceland. However, the highest O\textsubscript{3} concentrations at both stations were observed on 3 May 2017 when the air masses were transported from Europe (Fig. 10b). The air masses arrived in Svalbard from the west and did not pass over the areas with elevated BrO concentration.

4 Discussion

The NO\textsubscript{x} monitor in Adventdalen was located far away from stationary emission sources and showed the highest daytime NO\textsubscript{2}/NO\textsubscript{x} ratio (Table 1). We would like to investigate how the ratios observed there were affected by photolysis. The
Table 2. Median values of measured parameters and the three most often observed wind directions for different weather regimes and for the whole campaign. The concentrations exceeding median value for the whole campaign are shown with bold font.

| Parameter | 23 Mar–7 Apr: no AR | 7–9 Apr: ScTr | 9–16 Apr: AR | 16–24 Apr: AR | 24–26 Apr: ScTr | 26–29 Apr: no AR | 29 Apr–5 May: ScBL | 5–13 May: GL | 13–15 May: no AR | Whole campaign |
|-----------|---------------------|--------------|--------------|---------------|--------------|----------------|----------------|--------------|----------------|----------------|
| NO\textsubscript{x} (ppb): | Adventdalen | 0.44 | 0.42 | 0.54 | 0.31 | 0.55 | 0.36 | 0.36 | 0.22 | 0.26 | 0.35 |
| | Barentsburg | 0.01 | 0.38 | 0.02 | 0.01 | 0.01 | 0.44 | 0.06 | 0.01 | 0.06 | 0.01 |
| | Ny-Ålesund | 0.26 | 0.16 | 0.15 | 0.08 | 0.09 | 0.00 | 0.00 | 0.02 | 0.00 | 0.12 |
| O\textsubscript{3} (ppb): | Zeppelin | 37.76 | 36.28 | 43.29 | 38.83 | 37.28 | 41.69 | 33.57 | 32.07 | 39.84 | 37.68 |
| Wind speed (m s\textsuperscript{-1}): | Adventdalen | 3.9 | 7.0 | 4.9 | 7.8 | 4.1 | 4.6 | 3.3 | 4.6 | 3.2 | 4.6 |
| | Barentsburg | 3.2 | 1.4 | 1.7 | 1.2 | 1.0 | 3.3 | 2.1 | 2.3 | 2.7 | 2.2 |
| | Ny-Ålesund | 3.9 | 1.5 | 2.4 | 3.0 | 1.6 | 4.0 | 3.2 | 3.2 | 4.5 | 3.2 |
| Temp. at 2 m (°C): | Adventdalen | −15.1 | −6.6 | −8.9 | −12.4 | −9.9 | −6.8 | −10.6 | −11.4 | −8.8 | −9.9 |
| | Barentsburg | −14.5 | −6.5 | −6.8 | −8.4 | −8.4 | −6.5 | −8.6 | −11.4 | −6.8 | −9.9 |
| | Ny-Ålesund | −15.1 | −7.5 | −6.4 | −8.6 | −1 | −2 | −12.4 | −9.9 | −1 | −2 |
| PBL (m): | Adventdalen | 299.4 | 252.3 | 251.1 | 270.5 | 146.2 | 635.4 | 366.3 | 373.4 | 269.6 | 298.2 |
| | Barentsburg | 366.3 | 191.1 | 256.6 | 319.1 | 277.4 | 616.0 | 447.1 | 503.7 | 526.6 | 379.2 |
| | Ny-Ålesund | 182.9 | 54.2 | 155.3 | 150.1 | 117.7 | 501.8 | 243.2 | 208.0 | 677.8 | 185.7 |
| UV-A + UV-B (W m\textsuperscript{-2}): | Longyearbyen | 1.4 | 3.4 | 3.1 | 4.7 | 4.3 | 4.9 | 3.5 | 8.6 | 5.2 | 3.8 |
| | Ny-Ålesund | 2.4 | 6.1 | 5.1 | 7.9 | 5.4 | 7.5 | 7.7 | 13.1 | 7.1 | 6.3 |

Note: The concentrations exceeding median value for the whole campaign are shown with bold font.
Figure 8. Synoptic-scale meteorological conditions in ERA5 reanalysis data for the nine sub-periods. The colour scale and the white contour lines show air temperature at 2 m height and mean sea level pressure, respectively. The black arrows represent the prevailing wind direction and show the length relative to the wind speed. They are plotted with a resolution of 2° longitude and 1° latitude.

Figure 9. FLEXPART trajectory probability for 10 d backward trajectories (dark green contours with step of 0.001) and GOME2 BrO vertical column density (VCD) (colour scale) for the different sub-periods. The percentage of trajectories descending from higher altitudes (> 2000 m) is 7%, 12%, 27%, 9%, 9%, 33%, 18%, 4% and 24% for the sub-periods in (a)–(i), respectively.
photolysis rate of NO\(_2\) depends on solar zenith angle (Parrish et al., 1983), which in turn depends on day of year. Measurements were performed between days 81 and 134, and the noon solar zenith angle in Longyearbyen area varied from approximately 77 to 62° (Robertson et al., 2006). Following Eq. (15) in Parrish et al. (1983), the minimum clear-sky photolysis rate for the start of the campaign was 0.0026 s\(^{-1}\), and the maximum clear-sky photolysis rate for the end of the campaign was 0.0061 s\(^{-1}\) (black squares in Fig. A1a in Appendix A). There are many factors that affect NO\(_2\) photolysis rate, such as aerosol load, clouds, water vapour content and surface albedo (Trebs et al., 2009). The albedo may significantly increase the NO\(_2\) photolysis rate (Trebs et al., 2009), and Dickerson et al. (1982) suggested albedo of snow with respect to \(j(\text{NO}_2)\) to be 93%. Trebs et al. (2009) suggested in their Eq. (2) a polynomial fit between global irradiance and NO\(_2\) photolysis rate that includes both clear-sky and cloudy conditions and takes into account the contribution of albedo. The albedo calculated as the ratio of upward and downward short-wave radiation measured by a CNR1 net radiometer (Kipp & Zonen) in Adventdalen and observed global radiation were used to estimate \(j(\text{NO}_2)\) (red line in Fig. A1a). Figure (A1b) shows the NO/NO\(_2\) ratio calculated using O\(_3\) concentration measured in Barentsburg (closest station where O\(_3\) measurements were available), \(j(\text{NO}_2)\) and the temperature-dependent rate coefficient \(k_{\text{NO}_2+\text{O}_3}\) obtained using temperatures in Adventdalen (Eq. 6.6 and Table 6.1 in Seinfeld and Pandis, 2006). The peaks of NO/NO\(_2\) ratios are especially pronounced for the days with decreased O\(_3\) concentration (1 April 2017 and the period from 4 to 9 May 2017). Note that the calculation is based on the O\(_3\) data from Barentsburg; thus this introduces an uncertainty in the exact NO/NO\(_2\) ratios estimated for Adventdalen. The observed and calculated NO\(_2\)/NO\(_x\) ratios for Adventdalen are shown in Fig. (A1c). The missing data in the observed NO\(_2\)/NO\(_x\) ratio (light blue line) indicate that both NO and NO\(_x\) values were within zero-noise level, while missing data in the calculated NO\(_2\)/NO\(_x\) ratio are due to missing O\(_3\) observations in Barentsburg. The observed and calculated values are of the same order, but the NO\(_2\)/NO\(_x\) ratio is underestimated in 64% of all available data, especially for the days with low O\(_3\) values. This underestimation was present, even in hours influenced by fresh local NO emission (light purple line), and might have resulted from the modelling errors that could occur if the surface albedo was high (Trebs et al., 2009) or because the actual O\(_3\) values in Adventdalen were lower than in Barentsburg. The NO\(_2\)/NO\(_x\) ratio is overestimated in 31% of all available data. In these hours, the actual O\(_3\) concentration might have been higher in Adventdalen than in Barentsburg (used for calculations). The most pronounced overestimation is noticeable in the period from 26 to 29 April when NO values in Barentsburg were higher than in Adventdalen, and thus more pronounced O\(_3\) titration with local NO might have occurred in Barentsburg.

The results from radiosonde and ozone soundings as well as CO and particle measurements, presented in this study, demonstrate that the O\(_3\) observations at the Zeppelin station were not sensitive to the local NO\(_x\) pollution from Ny-Ålesund and thus were representative as background values for comparison with Barentsburg and investigation of the influence of prevailing long-range transport patterns on the measurements at these stations. Furthermore, O\(_3\) data from the Zeppelin station may be used to assess how the PAN decomposition might have affected the background NO\(_x\) concentrations in Svalbard during the 2017 campaign. Previous studies have shown that the NO\(_x\)/PAN ratio in-
creases at temperatures above \(-10^\circ\text{C}\), and PAN decomposition becomes a major source of background \(\text{NO}_x\) in Svalbard (Beine et al., 1997a; Beine and Krognes, 2000). As the temperature at the Zeppelin station varied from \(-22.7\) to \(0.8^\circ\text{C}\) during the campaign, we would like to investigate the contribution of PAN decomposition to the background \(\text{NO}_x\) concentration in Svalbard. The PAN decomposition rate may be estimated using several approaches (Beine et al., 1997a); here we apply a linear relationship between \(\text{O}_3\) and PAN concentration derived from previous measurements at the Zeppelin station, \(\text{PAN \ [ppt]} = (\text{O}_3 \ [\text{ppb}] - 26.58)/0.034\), and then we calculate the PAN decomposition rate (Beine and Krognes, 2000) (Fig. A2a). The maximum PAN decomposition rate has been calculated using temperatures and \(\text{O}_3\) concentration observed at the Zeppelin station, applying Eq. (1) from Beine and Krognes (2000). The depletion events when \(\text{O}_3\) concentration was below 26.58 ppb have been excluded from the calculation (Beine and Krognes, 2000). The median calculated PAN concentration of 356 pptv \((10^{-3}\ \text{ppb})\) is comparable with previous springtime Arctic observations (Beine and Krognes, 2000; Kramer et al., 2015). The estimated maximum PAN decomposition rate for the whole campaign varied from \(-0.0033\) to \(-17.2\ \text{pptv h}^{-1}\), with a median value of \(-1.29\ \text{pptv h}^{-1}\) (Fig. A2b). The maximum PAN concentration coincides with the strongest \(\text{O}_3\) increase event which occurred on 3 May 2017 (Fig. 10b). The temperature increased simultaneously for that day (Fig. 2a), promoting efficient PAN decomposition (Fig. A2b). Applying Theil’s non-parametric regression with a slope of \(-5.07\ \text{(pptv NO}_x\ \text{pptv}^{-1}\ \text{PAN}^{-1})\) suggested by Beine et al. (1997a) for Svalbard, the background concentration of \(\text{NO}_x\) would be 87.2 pptv. However, these concentrations are too low for the equipment used in the 2017 campaign to detect the variations in the concentrations caused by the PAN decomposition.

The absence of collocated \(\text{NO}_x/\text{O}_3\) measurements in Ny-Ålesund and Longyearbyen does not allow us to investigate how the local emissions affect \(\text{O}_3\) concentrations in these settlements. This is a drawback of this study. The \(\text{O}_3\) monitoring at the Zeppelin station is a long-term ongoing research project, and relocation of the instrument to the village from the mountain observatory would introduce bias in the long-term atmospheric composition observations. The study in Adventdalen was the first combined air pollution and meteorological fieldwork in Longyearbyen. The measurements there were done by the main author, and only the \(\text{NO}_x\) monitor was installed there due to the limited grant funding.

To investigate the influence of local \(\text{NO}_x\) emissions on the \(\text{O}_3\) concentration in Ny-Ålesund, as is required in the third hypothesis stated in the introduction of the current paper, we may use historical observations. The data from only six \(\text{O}_3\) sonde launches were available for the 2017 campaign (Fig. 4). However, the long-term data below 100 m from the \(\text{O}_3\) sonde profiles may be used to study influence of the local \(\text{NO}_x\) pollution in Ny-Ålesund on the \(\text{O}_3\) concentration.

These observations are suitable for this purpose because the \(\text{O}_3\) sonde launching facility is located just 200 m to the south-west and 500 m to the south from the \(\text{NO}_x\) monitor and diesel power plant, respectively. Thus, when the monitor detected \(\text{NO}_x\) concentration above the long-term springtime average in the launch hour, the influence of locally polluted air masses might have been observed in the lowest \(\text{O}_3\) sonde data. There were in total 59 \(\text{O}_3\) sonde launches for which \(\text{NO}_x\) monitor data were available in spring 2009, 2010, 2015, 2016, 2017 and 2018. The \(\text{O}_3\) profile data in the lowest 100 m have been extracted for all 59 launches and grouped according to the \(\text{NO}_x\) concentration detected by the monitor and wind direction in the \(\text{O}_3\) sonde profiles: (1) above mean \(\text{NO}_x\) concentration and northerly wind direction and (2) below or equal to mean \(\text{NO}_x\) concentration and northerly wind direction. The median and mean \(\text{O}_3\) values below 100 m in the group where the \(\text{NO}_x\) values were above \(\text{NO}_x\) mean were 11 % and 15 % lower, respectively, than for the second group with northerly winds but without elevated \(\text{NO}_x\) concentration. Thus, the \(\text{O}_3\) concentration in lowest 100 m downwind from the power plant in the settlement may be reduced significantly due to local \(\text{NO}_x\) emissions, but the frequency of such events is unknown in the absence of continuous \(\text{O}_3\) measurements in the village.

The \(\text{NO}_x\) concentrations depended strongly on the wind direction at the stations located in the vicinity of the stationary pollution sources, Ny-Ålesund and Barentsburg. In turn, the wind direction at all the sites depended on the synoptic-scale conditions but was modified locally due to different mechanical and thermodynamic processes controlling local circulations such as katabatic winds and topography-induced wind channelling specific for each location (Esaü and Reina, 2012; Maturilli et al., 2013). Remarkably, the westerly component of the wind at all stations only appears when the synoptic-scale westerly wind brought warm air from the North Atlantic to the Svalbard inland during transition to, and during part of, the large-scale ScBL regime. This reverses semi-permanent thermal flow from the glaciers towards the sea prevailing in Svalbard in spring.

Our analysis of the trajectory probability for different weather regimes showed that the elevated median \(\text{O}_3\) concentrations were observed for the sub-periods when the air masses arrived from the south, west or east. In contrast, the long-range-transported \(\text{O}_3\), brought by the air masses from the north, may be affected by the regional \(\text{O}_3\) depletion north of Svalbard where the elevated concentrations of total BrO VCDs were detected in the satellite data. Similarly, Koo et al. (2012) studied advection of the \(\text{O}_3\)-depleted air masses and found that these events were driven by local or short-range (1 d) transport from the nearby region. A recent study of Bougoudis et al. (2020) explored connection between first-year sea ice and bromine explosion events. In spring (March, April, May) 2017, Arctic mean tropospheric BrO VCDs over sea ice were on the order of \(4\times 10^{15}\) molec. cm\(^{-2}\), and significant anomalies of tropospheric BrO VCDs were observed.
over the sea ice north of Svalbard at approximately 85° N. In addition to the sea-ice conditions, the tropospheric BrO plume formation depends on various meteorological factors and the amount of blowing snow (Bougoudis et al., 2020). To investigate these processes, the weather regime approach presented in the current study may be applied together with the long-term BrO remote-sensing data and in situ measurements from the Arctic stations in further studies.

However, as we show in the analysis of the extreme O₃ depletion and increase episodes, the specific short-term events of long-range transport need to be investigated separately as there is a spread in air mass origin and transport altitude for the longer periods defined by the weather regime classification, especially for the “no regime” situation.

To get a more robust result linking weather regimes and air quality, we would like to compare long-term springtime (23 March–15 May) weather regime data with NOₓ data from Ny-Ålesund, O₃ concentration from the Zeppelin station, O₃ sonde and radiosonde data as well as FLEXPART trajectories for the period from 1990 to 2018. The FLEXPART and weather regime data were available for all years, while there were gaps in observational data from Ny-Ålesund. The data availability chart is shown in Fig. A3, indicating the number of hourly measurements for surface NOₓ and O₃ data and the number of radiosonde and ozone sonde launches per spring season each year. The hourly O₃ data are available for all years, while hourly NOₓ data were only available in 2009, 2010, 2015, 2016, 2017 and 2018. After spring 2018, the NOₓ monitor was moved to another location in Ny-Ålesund; therefore 2019–2022 data are not included in the current analysis to keep measurement consistency. The O₃ soundings and radiosonde AWT’s data sets start in 1992 and 1993, with the median number of radiosonde soundings and O₃ soundings per spring season being 54 and 11, respectively.

The box and whisker plots of O₃, NOₓ and temperature inversion strength (TIS) for different weather regimes are shown in Fig. A4. As during the 2017 campaign, the highest median O₃ values were detected for the ScTr regime, while the zonal regime (ZO), during which O₃ values were also higher, was absent during the fieldwork (Fig. A4a). The ZO regime is characterized by a negative geopotential height anomaly at 500 hPa centred between Iceland and the southern tip of Greenland and southerly flow over Svalbard (Papritz and Grams, 2018). The lowest median O₃ values in the long-term data are for GL regime and for European blocking (EUBL) that was absent in spring 2017. The variability of O₃ concentrations (range between the 25th percentile and 75th percentile) for the EUBL regime is also remarkably higher than for other regimes. The EUBL regime is characterized by the positive 500 hPa anomaly centred over the North Atlantic. This promotes transport of air from south-west and west to Svalbard (Papritz and Grams, 2018). Long-term trajectories for the different regimes are shown in Fig. A5. The lowest percentage of trajectories descending from higher altitude (> 2000 m) is modelled for GL regime, while the highest percentage of elevated trajectories is obtained for EUBL, ScBL and ZO regimes. No specific trajectory probability pattern may be defined for “no regime” conditions (Fig. A5a), while distinct long-range transport signatures are identified for the other seven regimes (Fig. A5b-h). In addition to the air transport path and trajectory altitudes, the sea-ice conditions and BrO concentration are important factors affecting the concentration of O₃ in each particular season.

Similar to the 2017 results, the highest median NOₓ values are observed for “no regime” and ScTr, but two other regimes, when the long-term median NOₓ values are high as well, EUBL and ZO, were not present during the campaign (Fig. A4b). The EUBL shows pronounced transport of air masses from the west to Svalbard (Fig. A5f). Thus, a westerly component of the wind at the measurement stations and significant changes in local pollution dispersion conditions are expected for this regime, as was observed during the ScBL regime in 2017, when a westerly component of the wind was present as well.

Temperature inversions are common phenomena at high latitudes, in particular during the cold seasons due to radiative cooling of the surface and descending motion and heat advection from the south aloft. The inversions were detected on 27 % of all the days in the measurement campaign period in 2017. This frequency of inversion occurrence is quite low in comparison with the results from previous studies of Dekhtyareva et al. (2018), where it was observed in 60 % of the springtime profiles in 2009. Despite low frequency of occurrence, temperature inversions have significant influence on the dispersion efficiency, and, hence, according to the WRS test, the median daytime (from 06:00 to 18:00 UTC) concentrations of NOₓ were higher ($p < 0.05$) at all three stations for the days when this phenomenon was observed in the radiosonde data in 2017. In the long-term data, the median temperature inversion strength was high for “no regime”, GL, AR and ScBL, but the highest median TIS was for the Atlantic trough (AT) regime, the regime that was absent during the campaign (Fig. A4c). The AT regime is characterized by a negative 500 hPa geopotential height anomaly to the east of Ireland and high cyclone frequency in that region (Papritz and Grams, 2018), while the cyclonic activity around Svalbard is lower, and these conditions may promote strengthening of the temperature inversion.

Thus, the results of the weather regime analysis performed for the 2017 campaign are representative of the characterization of the influence of different synoptic-scale conditions on the NOₓ and O₃ concentration in Ny-Ålesund. However, the three regimes that were absent during the 2017 campaign (AT, EUBL and ZO) are important in the long-term statistics for NOₓ, O₃ and TIS in the settlement.
5 Conclusions

Despite decades of industrial activity in Svalbard, there is no continuous air pollution monitoring in the region’s settlements except Ny-Ålesund. The NO\textsubscript{x} measurement results from the three-station network, Ny-Ålesund, Barentsburg and Longyearbyen, and O\textsubscript{3} data from Ny-Ålesund and Barentsburg have been compared for the first time.

A diurnal pattern in concentration of NO\textsubscript{x} at all three stations has been observed and attributed to variable emissions from the local sources of NO\textsubscript{x}. However, only data from Barentsburg and Adventdalen station show a significant change in the NO\textsubscript{2}/NO\textsubscript{x} ratio during the day, since the station in Ny-Ålesund is located close to a diesel power plant, a stationary source of fresh NO\textsubscript{x} emissions contributing to higher NO concentration. The NO\textsubscript{2}/NO\textsubscript{x} ratio observed in Adventdalen is comparable with modelling results obtained using radiation data from the valley and O\textsubscript{3} concentration from Barentsburg. Local emissions of NO\textsubscript{x} in Barentsburg may reduce O\textsubscript{3} concentrations in the settlement by a few percent from the background value due to O\textsubscript{3} titration. As has been shown from the analysis of the long-term O\textsubscript{3} sonde record, NO\textsubscript{x} emissions in Ny-Ålesund may affect O\textsubscript{3} concentration in the lowest 100 m downwind, but no influence of local pollution has been detected at the Zeppelin station at 474 m a.s.l. in spring 2017. There was no statistically significant difference in daytime and night-time O\textsubscript{3} values measured in Barentsburg and at the Zeppelin station, and both sites showed similar O\textsubscript{3} concentration dynamics controlled by long-range air mass transport.

The weather regime approach is novel in Svalbard air pollution research. This method has been used in the current study to identify the influence of large-scale circulation on local and long-range-transported air pollution in the area with complex topography. As expected, the large-scale wind is channelled by the local topographical features, and this determines the wind direction and speed in all three settlements, and therefore the correlations of NO\textsubscript{x} concentrations between the stations are weak. In Ny-Ålesund and Barentsburg, the stations are located so that downwind concentrations from the local sources are observed rarely, since the prevailing wind direction is different. The measurements in Adventdalen have been made downwind from the source, since both the snowmobile route and prevailing wind direction are along the valley. However, traffic is a temporary source of emissions, and the mean wind speed in Adventdalen valley is high, and therefore mean NO\textsubscript{x} concentrations there are low. Despite low correlation between the NO\textsubscript{x} values from the three stations, there are common synoptic conditions that promote accumulation of local pollution in the settlements, namely low wind speed and air temperature and the presence of temperature inversions. In contrast to NO\textsubscript{x}, the concentrations of O\textsubscript{3} in Barentsburg and at the Zeppelin observatory are moderately correlated and depend on synoptic conditions that promote transport of air masses enriched or depleted in O\textsubscript{3}. In other words, both these stations are regionally representative of the O\textsubscript{3} concentrations.

The large-scale weather regimes control the synoptic meteorological conditions and determine the atmospheric stability and efficiency of local pollution dispersion. The analysis of the long-term weather regime, trajectory and observational data from Ny-Ålesund supports our findings from the 2017 campaign. The lowest median O\textsubscript{3} values were identified for the Greenland blocking regime, when the trajectories were arriving more frequently from the sea-ice-covered regions, and the percentage of high-altitude trajectories that might bring O\textsubscript{3}-enriched air was low. The highest median O\textsubscript{3} values were observed for the Scandinavian trough regime, characterized by the relatively high percentage of high-altitude trajectories and trajectories arriving from south-east of Svalbard. During this regime, NO\textsubscript{x} concentrations were elevated as well. The highest temperature inversion strength was observed for “no regime”, Greenland blocking, Atlantic ridge and Scandinavian blocking. The maximum background NO\textsubscript{2} concentration originating from PAN decomposition was modelled for the strongest O\textsubscript{3} increase event that occurred during the Scandinavian blocking regime. At the same time, three regimes absent during the 2017 campaign (Atlantic trough, European blocking and Zonal regime) appeared to be significant for NO\textsubscript{x}, O\textsubscript{3} and temperature inversion statistics in Ny-Ålesund. However, the effect of each weather regime on the air quality in different settlements depends on the local features such as pollution sources and wind channelling; thus it would be of interest to compare long-term NO\textsubscript{x} and O\textsubscript{3} data from Barentsburg with weather regime data in further studies.

The application of the weather regime approach in air quality study for the three Svalbard stations allows us to facilitate prediction of the conditions promoting long-range transport to and accumulation of local pollutants at the measurement sites. The weather regimes typically persist for a period of 10 d and longer. Hence, joint intensive observational campaigns may be planned ahead at any of the three stations depending on the expected conditions. This provides a new opportunity for the collaboration in atmospheric research in Svalbard and allows more effective organization of specific field observations devoted to, for example, the study of photochemical reactions in the polar atmosphere, investigations of the influence of turbulence and stability on air pollutant dispersion, and studies on aerosol and cloud interaction.
Appendix A

Figure A1. (a) $j_{NO_2}$ calculated as a function of solar zenith angle in clear-sky conditions (Parrish et al., 1983) (decreasing from 77 to 59° for solar noon time in Adventdalen from 23 March to 15 May 2017 (Robertson et al., 2006)) and as a function of observed global radiation and albedo (Trebs et al., 2009); (b) NO/NO$_2$ ratio calculated using O$_3$ concentration measured in Barentsburg, $j_{NO_2}$ and temperature-dependent rate coefficient $k_{NO_2+O_3}$ obtained using temperatures in Adventdalen (Eq. 6.6 and Table 6.1 in Seinfeld and Pandis, 2006); (c) observed and calculated NO$_2$/NO$_x$ ratio for Adventdalen.

Figure A2. (a) PAN concentration calculated using the linear relationship between PAN and O$_3$ concentration suggested by Beine and Krognes (2000). (b) PAN decomposition rate calculated using Eqs. (1) and (2) from Beine and Krognes (2000).
Figure A3. The O$_3$ and NO$_x$ hourly observations and O$_3$ sonde and radiosonde data availability chart for spring seasons 1990–2018.

Figure A4. (a) O$_3$ concentrations for different weather regimes for spring seasons 1990–2018. (b) NO$_x$ concentrations for different weather regimes for spring seasons 1990–2018. (c) Temperature inversion strength for different weather regimes for spring seasons 1990–2018. The maximum values that exceed y-axis limits in plots (b) and (c) are shown with red stars.
Figure A5. FLEXPART trajectory probability for 10 d backward trajectories for the different weather regimes for spring seasons 1990–2018. The percentage of trajectories descending from higher altitudes (> 2000 m) is 18 %, 14 %, 13 %, 15 %, 17 %, 22 %, 21 % and 21 % for the regimes in plots (a)–(h), respectively.

Data availability. The radiosonde data for March 2017 and for April–May 2017 are available via the GRUAN home page (https://doi.org/10.5676/GRUAN/RS92-GDP.2, Sommer et al., 2012) and in the database PANGAEA (https://doi.org/10.1594/PANGAEA.879767, Maturilli, 2017a and https://doi.org/10.1594/PANGAEA.879820, Maturilli, 2017b), respectively. The analysed O₃ sonde data are stored in the Network for the Detection of Atmospheric Composition Change (NDACC) archive at https://www-air.larc.nasa.gov/missions/ndacc/data.html?station=ny.alesund/ames/o3sonde/ (von der Gathen and Rex, 2020). The NOₓ data from Adventdalen are available at the UiT Open Research data portal (https://doi.org/10.18710/TXQ7EV, Dekhtyareva, 2018).

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