We thank the reviewer for his/her comments that have improved the completeness and clarity of this manuscript. The comments by the reviewer are given below in black, our responses are marked in red and the corresponding changes to the manuscript are in *italics*.

Interactive comment on “Characterization of Transport Regimes and the Polar Dome during Arctic Spring and Summer using in-situ Aircraft Measurements” by Heiko Bozem et al.

Anonymous Referee #1

Received and published: 6 March 2019

This manuscript provides an analysis of air masses present during the 2014 and 2015 NETCARE airborne measurements spanning a broad region of the western Arctic between Spitsbergen and Alaska. An overview of the meteorological conditions during the summer (2014) and winter/spring (2015) campaigns is given, then trajectories are used to identify airmass history. Trace gas observations are described, then combined with potential temperature to identify regions with sharp gradients in CO and CO\textsubscript{2}. These gradients are used to define the "polar dome", the region of the cold, stable, near-surface Arctic airmass that is most isolated from midlatitude influences. (As this airmass wobbles over sources of pollution in the winter, it accumulates pollutants because sinks are very slow, leading to the seasonal near-surface "Arctic haze" phenomenon, which is of broad interest.)

The statistics of CO and CO\textsubscript{2} abundance in the different regions identified from this analysis are then presented. Next, a "transport regime" analysis, based on the trajectories and using methods developed in Binder (2017), is used to evaluate the influence of lifting within and outside the Arctic, and diabatic processes, on CO and CO\textsubscript{2} mixing ratios within and outside of the polar dome. Finally, a discussion section evaluates the abundance of nucleation and Aitken mode particles within the three regimes identified in the earlier analysis (inside the polar dome, outside the polar dome, and a mixing region).

This is an ambitious manuscript, with many parts. It has interesting sections, but it doesn’t seem to have a strong overall purpose. My fundamental complaint with the manuscript is that it doesn’t make the case for any generality to the analysis. Are the results more broadly applicable outside of the narrow time period and location of the NETCARE airborne observations in 2014 and 2015? For example, in Sect. 5.3 there is a long discussion of how the CO and CO\textsubscript{2} observations can be used to identify the polar dome boundaries, and a specific range of potential temperatures and latitudes is the result of the analysis. This is great for these NETCARE observations, but are these findings more broadly applicable? For example, could one take the long-term surface observations at UtqiaÁaÁvik (Barrow) or Alert or Zeppelin, apply the potential temperature and CO/CO\textsubscript{2} screens developed in this manuscript, and separate the data out into "in the polar dome" and "out of the polar dome" datasets? This would be useful to the scientific community. Without such broader relevance, this analysis is of interest only to the very small set of scientists interested in the NETCARE data.

We appreciate the reviewer’s comments and the valuable suggestions for restructuring the manuscript which we largely followed and extended the discussion of the transport regimes. Regarding the point of a larger scale applicability of the method we included a first comparison with station data from Zeppelin in the reply.
In this particular paper our intention was indeed to develop an empirical dome boundary based on the airborne measurements of tracer like CO and CO$_2$ for NETCARE and to show first applications. We will extend the method and apply it to other airborne data sets or ground based data. Note however, that ground based data could eventually have a different tracer characteristics since these data are stronger affected by local sources and sinks and partly decoupled from the free troposphere by the boundary layer inversion. Nonetheless, we tested the approach for Zeppelin, which provides very promising results (see below). We will publish a follow-up paper, which will apply the method to a larger data set to investigate the general applicability.

In addition to my concern with the applicability of the findings, I feel the manuscript also needs restructuring. The paper opens with the meteorological analysis, which is fine. Next, though, is the trajectory analysis. It would be more logical if the next section were the identification of the polar dome using the trace gas gradients. Then the backtrajectories could follow, with backtrajectories initiated either within the polar dome, outside of it, or in the mixing region. The trajectories would then provide an independent and intuitive confirmation of the identification of the polar dome that was derived from the trace gas and potential temperature data.

We restructured the manuscript mostly as suggested: We moved the detailed trajectory analysis of former Figure 4 and 5 to the supplement and combined the chemical regime discussion based on trace gases with the (newly restructured) aerosol paragraph at the end. This results in a structure as suggested: Meteorological analysis, determination of the polar dome boundary, transport regimes and air mass history, chemical regimes. Note that Willis et al. (2019) and Schulz et al. (2019) already used the polar dome boundary presented in this study to analyse aerosol composition, transport and sources within the springtime polar dome and within and partly outside the summertime polar dome, respectively. For a more detailed aerosol analysis we refer to their studies and only briefly address characteristics of aerosol within this study.

Section 5.4, which is a presentation of PDFs of CO and CO$_2$ from the three different air masses is not very logical. CO and CO$_2$ were used to identify the three air mass classifications, after all, so it’s entirely expected that they would have different PDFs—the reasoning is circular. It is important to keep in mind that the dome boundaries were determined based on the gradients of CO and CO$_2$, not applying any threshold value for tracers. In the case of horizontal (i.e. isentropic) gradients the analysis was performed for different isentropes in intervals of 2K, allowing for different thresholds for each isentropic interval. There is no step within the analysis, which makes use of an absolute value of either CO or CO$_2$. Since the gradient is not related to an absolute value of CO or CO$_2$, trace gas values and particular the variability therefore still provide additional information for the regimes. The width of the distributions should be larger outside the dome regions, the mean (median values) resemble the surface network data. We therefore kept this section since it illustrates the chemical properties of the regimes. As mentioned before it is moved to the end of the manuscript.

Instead of this section, the next logical section would be examining the transport regimes using the Binder et al. methodology, as it continues the analysis of trajectories. After restructuring the paper, the next section that follows is the analysis of the transport regime based on the Binder et al. methodology as suggested by the reviewer.

This section could be made stronger by coupling the delta-theta/delta-T plots with graphs of trajectory clusters (e.g., plots trajectory altitudes as a function of latitude). This would bolster the rather speculative discussion about the meaning of each of the sectors of the Binder plots. This section could get rather long, so it might make sense to give one example (from the springtime flights, perhaps) and place the rest in the supporting materials. This analysis is the part of the current manuscript that is really informative outside of the narrow range of the NETCARE project, as it suggest broader generalities about how transport occurs into the polar dome. I’d like to see it...
developed more thoroughly and the conclusions made firmer, perhaps with a concluding paragraph that summarizes the findings from this section.

We included the figures according to the suggestion showing the evolution of the trajectories. As an example, sector 1 of the April 2015 measurements was chosen for those data points inside the polar dome, since it is the most dominant sector. In contrast, sector 4 was chosen for the air masses outside the dome, representing the largest contribution there. For better figure clarity only every 20th individual trajectory is plotted. Corresponding figures for the dominating sectors for July 2014 are included in the supplement material.

New figures for trajectory analysis inside the polar dome. Shown is the dominating sector 1 from Fig. 11b in the paper:

![Figure 12: (a) Trajectories of the most dominant sector 1 for air masses inside the polar dome. The color code represents the pressure along the trajectories. (b) The same trajectories as in (a) as a function of pressure and latitude, color coded by potential temperature. In both figures (a) and (b) black circles denote the initialisation point of the trajectory along the flight track. The black open squares show the position of the trajectory 10 days back in time. Figures (c) and (d) show a vertical cross section of the trajectory evolution over the 10 days of travel with the color code denoting the temperature (c) and potential temperature (d). The black line marks the median pressure of the trajectory cluster at the individual time steps and the grey line indicates the median temperature and median potential temperature, respectively. Note that in all figures only every 20th trajectory is plotted for figure clarity.](image-url)
New figures for trajectory analysis outside the polar dome showing the dominating sector 4 from Fig. 11c in the paper:

Figure 13: (a) Trajectories of the most dominant sector 4 for air masses outside the polar dome. The color code represents the pressure along the trajectories. (b) The same trajectories as in (a) as a function of pressure and latitude, color coded by potential temperature. In both figures (a) and (b) black circles denote the initialisation point of the trajectory along the flight track. The black open squares show the position of the trajectory 10 days back in time. Figures (c) and (d) show a vertical cross section of the trajectory evolution over the 10 days of travel with the color code denoting the temperature (c) and potential temperature (d). The black line marks the median pressure of the trajectory cluster at the individual time steps and the grey line indicates the median temperature and median potential temperature, respectively. Note that in all figures only every 20th trajectory is plotted for figure clarity.

The following paragraph was added to the manuscript:
Based on the results from the phase space diagrams we further analyze the trajectories of the individual clusters. This allows for a more detailed analysis of the physical processes along the trajectory. Therefore, we compare the two most dominant sectors for the April 2015 measurements in Figs. 12a-d and 13a-d. Sector 1 is mostly dominated by air masses confined to the central Arctic at all altitude levels (see Figs. 12a and b). The air masses show a weak descent during the 10 days before the measurements but experience a very pronounced decrease in temperature and potential temperature indicated by the evolution of median temperature of the whole trajectory cluster (see Figs. 12c and d). In contrast, the trajectory analysis of the most dominant sector 4 of air masses
outside the polar dome reveals different air streams contributing to the cluster. Air masses originate at different altitudes in the central Arctic, at low level over the Pacific Ocean and from the upper troposphere over Asia. Air masses in this cluster are characterized by a significant increase of median temperature and decrease of median potential temperature indicating a descending trend, which is confirmed by the median pressure decrease over the time of travel. The low-level transport over the Pacific is associated with a low-pressure system over Alaska. Those air masses arrive at the polar dome boundary in the measurement region after experiencing a week net cooling over Alaska.

We conclude that air masses within the aged summertime polar dome are mostly confined to the boundary layer while they experienced a week diabatic warming due to insolation in July 2014 during NETCARE. In the mixing region and outside the polar dome diabatic cooling and a continuous descent is observed. Within the polar dome in April 2015 during NETCARE mostly near-surface processes (diabatic cooling due to the flow over cold surfaces) dominate the recent transport history of air masses in the lower polar dome. Air masses in the upper polar dome experience a very slow descent induced by radiative cooling. Outside the polar dome air masses mostly arrive at higher potential temperatures in the Arctic and experience a continuous slow descent with increasing temperatures but only week diabatic cooling.

Following the transport regime analysis, the next logical section would be to see how these different airmass types are manifested in the pollution loadings. This is currently Sect. 6, which is labelled “discussion”. I appreciate the novelty of Fig. 15, but it’s very difficult to understand the grey-scale coloring on top of the colored classification scheme. I’d much rather see PDFs of the number concentrations in the 5-20 and 20-100 and >100 size class. Are there also observations of aerosol composition (e.g., BC abundance, composition) that could be added to this section? You’ve effectively classified the measurements into airmass type; it would be extremely interesting to see how all the available aerosol microphysical and chemical parameters vary within these different airmasses, and compare them with existing literature values.

We thank the reviewer for this point and revised the paragraph in the revised manuscript. We just want to show the relevance of our empirical dome boundary to other applications using aerosol formation as an example. The specific analysis and discussion of aerosol processes is beyond the scope of this paper. We refer to Willis et al. (2019) and Schulz et al. (2019) for a more detailed discussion on aerosol composition within and outside the polar dome. They used the polar dome boundary derived in this study for their analysis. We revised the paragraph on aerosol as follows including a new figure 15:
Figure 15: Normalized probability density functions for particles with diameters between 5 and 20 nm (a) and larger than 100 nm for July 2014 (b). The colour code represents the three different regions identified during the polar dome analysis. Panel (c) shows the distribution of the fraction of particles containing trimethylamine (TMA) with respect to the total amount of particles measured by single particle mass spectrometry (Köllner et al., 2017) in the maximum potential temperature and median latitude coordinate system. Note that for the reason of particle spectra statistics the resolution of median latitude and maximum potential temperature interval is reduced compared to the refractory black carbon distribution (BC) shown in panel (d). The enhancement in BC within the polar dome is most probably due to fresh local pollution.

Having defined the polar dome based on trace gas gradients now allows for a more detailed study on aerosol associated with the different air masses. Efficient wet removal and less efficient transport from lower latitudes lead to generally low aerosol concentrations (Stohl, 2006; Engvall et al., 2008), especially within the Arctic lower troposphere during summer. This is consistent with results in Fig. 15a and b. Observations of elevated levels of accumulation mode particles ($N_{>100}$) can be associated with regions outside the polar dome and subsequent transport to the measurement region. In parallel, regions within the polar dome were characterized by $N_{>100}$ smaller than 100 cm$^{-3}$. In contrary, number concentrations of ultrafine particles ($N_{5-20}$) showed occasionally larger values within the polar dome compared to outside (Fig. 15a), indicating the formation of ultrafine particles occurred within the polar dome region (Burkart et al., 2017). Exemplary for aerosol composition, particulate trimethylamine (measured by single particle mass spectrometry) can be associated with sources within the polar dome (Fig. 15c), consistent with results in Köllner et al. (2017) and Willis et al. (2017). In contrast, the abundance of refractory black carbon can be linked to pollution sources outside the polar dome and subsequent transport to the measurement region (Fig. 15d; Schulz et al., 2019). To conclude, the method introduced in this study is a useful tool to combine Arctic aerosol observations with transport processes and sources within and outside the polar dome region.

Finally, if it’s feasible it would be wonderful to extend this analysis to the surface data from the long-term monitoring sites. This would show that the classifications developed here are more broadly applicable. At least an evaluation of whether the approach here is applicable to other cases, or is specific to NETCARE, is needed.

In a follow-up paper to this study we will extend our approach to a variety of different field campaigns in the Arctic covering different seasons and different locations. To demonstrate the applicability of the tracer based diagnostics to a broader data set we analyzed two examples of hourly ground based observations at the Zeppelin Mountain observatory (Ny Alesund, Spitsbergen) (see Fig. R2). Based on CO-CO$_2$-scatter plots the signatures and characteristics of both species inside the polar dome during the campaign phases are also found in the respective ground based observations. Note that ground based and airborne observations are in principle affected by different
processes and not necessarily linked. Particularly, the CO and CO$_2$ data at higher potential temperatures during the airborne campaigns are linked to higher altitudes as evident in both Figs. R1 a and b.

![Figure R1 (a): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) for July 2014. (b): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) for April 2015. The color code denotes potential temperature.](image)

Note however, the good agreement at low potential temperatures particularly for the April 2015 case with exactly the same slope and similar absolute values at the station as deduced from the aircraft data as inside dome (see Figs R2 a and b). Based on this analysis we conclude that the Zeppelin observatory was inside the polar dome for April 2014 and 2015 and most probably inside the polar dome with episodes outside the polar dome in July 2014 and 2015. The latter is indicated by the correlation following the characteristics of the mixing region (see Fig. R2a). However, potential temperatures were as low as those observed during the research flights (not shown).
In addition to these larger, structural issues, the manuscript needs some technical correction. The primary author is not a native English speaker, but English-speaking co-authors should step up and give the manuscript a round of thorough copy-editing. Verb tense is not used consistently, which is distracting and sometimes confusing. (Example, p. 17 lines 18-21 go from "now calculated" to "determine" to "finally used"). There are quite a few typos that a spell checker should find, and terms like "surfacenear" instead of "near-surface" are present. This manuscript has a lot of good analysis using an interesting and unique dataset. The CO and CO2 measurements look spot-on with the long-term ground monitoring network data, which is very encouraging since these airborne measurements can be tricky given the large background. With the suggested restructuring and a tighter focus on how the findings are more broadly applicable, the manuscript should be quite suitable to publish in ACP.

We thank the reviewer for this point. Copy-editing for the manuscript will be done by native English speakers.


Acknowledgement:

All atmospheric data from Zeppelin are publicly available in the EBAS database (http://ebas.nilu.no) and we thank Cathrine Lund Myhre and NILU - Norwegian Institute for Air Research for making the CO and CO2 observations from Zeppelin available.
We thank the reviewer for his/her comments that have improved the completeness and clarity of this manuscript. The comments by the reviewer are given below in black, our responses are marked in red and the corresponding changes to the manuscript are in italics.

**Interactive comment on “Characterization of Transport Regimes and the Polar Dome during Arctic Spring and Summer using in-situ Aircraft Measurements” by Heiko Bozem et al.**

**Anonymous Referee #2**

Received and published: 16 March 2019

This work makes use of the atmospheric tracers CO and CO2 measured on two NETCARE flight campaigns together with 10-day back trajectories to describe air mass transport into the high Arctic during spring and summer. The authors find distinct transition zones between the mid-latitudes and the polar dome for the two seasons based on tracer gradients. The tracer derived polar dome boundaries are subsequently applied to aerosol number concentration data. In addition, the authors explore different transport pathways of air masses into the Arctic using a previously published phasespace approach that relates the maximum change in potential and absolute temperature along the trajectory.

This manuscript is very comprehensive and represents a novel approach to localize the polar dome boundary and to characterize air mass transport into the Arctic. The work shall definitely be published with revisions as described below.

General remarks:

To give the findings more relevance for Arctic atmospheric research it would be of high interest, whether the derived polar dome boundaries are representative for other Arctic sectors and years as well? Or would each field campaign have to do their own analysis following this example? So a discussion on how far the results can be generalized is needed.

This study was intended to derive a tracer based diagnostic for the determination of the polar dome boundary. In a follow-up paper to this study we will discuss the application of the metrics to a comprehensive data set consisting of several field campaigns in the Arctic covering different seasons and different locations.

To demonstrate the general applicability of our campaign-based approach we analyzed two years of hourly ground based observations at the Zeppelin Mountain observatory (Ny Alesund, Spitsbergen) to show a more generalization of the tracer-based diagnostic (see Figs. R2 a and b). According to our campaign-based findings using the CO-CO$_2$ relationship the analysis of the ground-based data confirm the signatures and characteristics of both species inside the polar dome during the campaign. This potentially allows for a determination of the station location relative to the polar dome. Note, however, that ground based observations are in principle strongly affected by local sources and sinks. Depending on e.g. the strength of the boundary layer inversion these effects decouple the free troposphere above from these observations within the planetary boundary layer below.
Figure R1 (a): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) for July 2014. (b): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) for April 2015. The color code denotes the potential temperature.

Figure R2 (a): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) within the aged polar dome (blue), the mixing region (green) and outside the polar dome (red) for July 2014. (b): Tracer-tracer scatter plot of all aircraft data points (background + pollution plumes) within (blue) and outside (red) the polar dome for April 2015. To separate the different regions the tracer derived polar dome boundaries are used. Boundary values for each region are summarized in Tab. 2 in the paper. The black circles denote the ground based data from the Zeppelin mountain observatory (Ny Alesund, Spitzbergen) for the months July (a) and April (b) 2014 and 2015.

Note however, the good agreement at low potential temperatures particularly for the April 2015 case with exactly the same slope and similar absolute values at the station as deduced from the aircraft data as inside dome (see Figs R2a and b). Based on this analysis we conclude that the Zeppelin observatory was inside the polar dome for April 2014 and 2015 and most probably inside the polar dome with episodes outside the polar dome in July 2014 and 2015. The latter is indicated by the correlation following the characteristics of the mixing region (see Fig. R2a). However, potential temperatures were as low as those observed during the research flights (not shown).
There is no discussion on the uncertainties of the variables along the trajectory such as potential and absolute temperature. Particularly the vertical location of the polar dome boundary would be subject to the uncertainty. At least some discussion on how the ECMWF input data compares to the in-situ measurements should be added.

The figures below show a comparison of temperature based on in-situ data and temperature derived from the analysis data interpolated to the time and location of the measurement. Lagranto backward trajectories were initialised every 10s along the flight track based on GPS horizontal coordinates and pressure. Therefore, the variability of potential temperature is driven by the temperature difference between observation and analysis. To quantify this temperature difference, we analysed the temperature for July 2014 and April 2015. For the measurements in July 2014 the median difference between in-situ temperature data and analysis data is 0.31 °C (interquartile range: -0.71 - 1.72°C). For the April 2015 measurements, the respective median difference is 1.50°C (interquartile range: 0.69 - 2.14 °C). Thus, we conclude, that the local variability of the dome boundary is largely driven by the variability of tracer gradients and the uncertainty of transport history backward in time. The latter, however, we try to evaluate (at least qualitatively) by using median latitude and maximum potential temperature. Linking these to our trace gas observations should in turn resemble the latitudinal and vertical gradient and thus account for transport history.

Figure R3: (a): Scatter plot of in-situ ambient temperature measured on the aircraft and temperature from the analysis data set at the time of the initialisation of the trajectories along the flight track for July 2014. (b): Probability density function of the difference between in-situ and analysis temperature data set. (c): Box and whisker plot for the difference between in-situ and analysis temperature data set.

Figure R4: (a): Scatter plot of in-situ ambient temperature measured on the aircraft and temperature from the analysis data set at the time of the initialisation of the trajectories along the flight track for April 2015. (b): Probability density function of the difference between in-situ and analysis temperature data set. (c): Box and whisker plot for the difference between in-situ and analysis temperature data set.
We added the following to paragraph 3.3:

As a measure for the uncertainty of the temperature along the trajectory we calculated the median difference between temperatures measured in-situ on the aircraft and the corresponding temperatures interpolated to the initialisation point of the trajectory along the flight track based on analysis data. For the measurements in July 2014 the median is 0.31 °C (interquartile range: -0.71 - 1.72°C). For the April 2015 measurements, the respective median difference is 1.50°C (interquartile range: 0.69 - 2.14 °C)

The application of the polar dome boundary to aerosol data is intriguing and it would be very interesting to know how other aerosol properties relate to the boundary. Adding such information to the manuscript would make it even longer than it already is. I would recommend exploring whether a “part 2” manuscript on aerosols could make sense. The manuscript is partly repetitive and the abstract reads almost like an introduction. Both the abstract and the whole manuscript should be shortened. There are some recommendations in the attached PDF.

We thank the reviewer for this point and refer to a later comment.

Specific comments:

p. 7, l. 15: What is meant by “The stability of the instrument. . .”? Is this the accuracy? I wonder whether none of the instruments has been described before and whether such an extensive description here is necessary?

The instrument is regularly calibrated during the flights to check for longer term drifts. We use these drifts as a measure for the reproducibility, which we term stability. Note that post processing accounts for these slow drifts assuming a linear drift between the calibrations. Accuracy is further affected by the uncertainty of the calibration standards. For clarification, we added the following sentence:

“Stability is a measure for reproducibility and based on the mean drift between two subsequent calibrations which were performed during flights”.

The instrument and measurement principle of the Aerolaser ultra-fast carbon monoxide (CO) monitor model AL 5002 were described extensively in for example two papers by Gerbig et al. (1999) and Scharffe et al. (2013). The Licor LI-7200 closed CO$_2$/H$_2$O analyzer from LI-COR Biosciences GmbH, modified for airborne use, was only briefly described in earlier studies in the Arctic in the framework of the NETCARE project. We follow the suggestion of the reviewer and shorten the description of the carbon monoxide monitor only leaving the specification of uncertainties for this study. We kept the more extensive description of the CO$_2$ monitor since it is the first time of a more comprehensive description of the current setup.

p. 8, l. 19: What is a “very stable stratification” compared to a stable stratification? Is the ECMWF data the analysis or re-analysis data?

The term “stable stratification” is appropriate and the text is changed accordingly. ECMWF data is operational analysis data.

Figure 6: The scales can be enlarged in both panels: 200 for a) and 415 for b). This way, more details would be visible. The legends will find place somewhere else... What is the purpose of the figure except discussing the seasonal cycle? If a direct comparison of the NETCARE data with the stations is the goal, there should be zooms for the short periods of time. Currently, one cannot see much because the symbols are so large and cover everything. Why is Mace Head chosen as a reference?

The purpose of the figure is to show both the seasonal cycle of both species CO and CO$_2$ in the Arctic as well as in the mid-latitudes and to illustrate the latitudinal gradient between the Arctic stations and mid-latitude stations. For figure clarity, only Mace Head is shown representing a mid-latitude station. It was chosen because it is a GAW station with CO and CO$_2$ data available at the respective
time period and Mace Head is furthermore located at one of the entry routes of mid-latitude air masses that frequently enter the Arctic. The comparison between ground based observations and airborne data is not the prime purpose of the figure. Therefore, we show a more extended time period and not only a zoom on the period of the measurement campaign. Figures 6 (a) and (b) were modified for figure clarity and are now Figs 4 (a) and (b) after restructuring the manuscript.

Figure 4. CO (a) and CO$_2$ (b) seasonal cycle based on NOAA ground based measurements in Alert (Canada), Barrow (Alaska) and Mace Head (Ireland) for the years 2014 and 2015. Running means are shown for the respective station data (symbols). Mean aircraft data for altitudes < 200 m for individual flights are overlaid. Error bars (yellow and orange shading) for the aircraft data are too small to be visible. NETCARE 2014 data are in yellow and NETCARE 2015 data are in orange.

Figure 8 b: Why does the potential temperature increase below which CO maximum concentrations occur with decreasing latitude? Some explanation is needed. The measurements at lower latitudes were all performed from Inuvik. At that time generally higher temperatures were observed in Inuvik compared to the Arctic at similar altitude levels. Furthermore, the CO maxima in the two respective regions have different causes. In the Arctic lower troposphere high CO values are observed at low potential temperatures inside the polar dome as remnants of the wintertime maximum of CO due to the weak photochemical activity. In contrast enhanced CO values at lower latitudes are mainly caused by recent emissions (1-2 weeks old) potentially transported from Asian source regions into the measurement region by long-range transport. Since potential temperature at the surface increases in general with decreasing latitude (due to increasing temperature, see Fig. 5 in the paper) the CO maxima originating from lower latitudes affect higher isentropes. In addition, warm conveyor belt (WCB) type transport may have occurred leading to diabatic transport of CO to even higher altitudes and higher potential temperatures.

p. 28, l. 6-18: This paragraph is a result and should be moved to the results section instead of being added to the discussions. The paragraph on aerosol was located in the discussion section since it was intended to illustrate the relation of our empirically derived dome boundary when investigating e.g. aerosol processes. We followed the reviewer’s suggestion and moved the paragraph to the results section and revised the whole paragraph including figures.

p. 28, l. 10-13: How can you infer that secondary aerosol formation is responsible for the concentration difference in the blue area? Based on the information provided, those particles are either between 5 and 20 nm or > 100 nm. For the first option, I find it difficult to believe that there is evidence to relate the increase in 5 to 20 nm particles to secondary aerosol formation based on the AMS and ALABAMA measurements in Willis et al. (2017). These instruments do not cover the
relevant size range. If the second option is true, the information on SOA contributing to particle growth to form CCN does not make sense, because particles are already in the CCN size range, even for low supersaturations. What is the explanation for the difference between panel a and b? And what is the relevance for CCN? Please revise the statement.

We thank the reviewer for this point and revised the paragraph in the revised manuscript. We just want to show the relevance of our empirical dome boundary for other applications using aerosol formation as an example. The specific analysis and discussion of aerosol processes is beyond the scope of this paper. We modified the paragraph as follows:

Figure 15: Normalized probability density functions for particles with diameters between 5 and 20 nm (a) and larger than 100 nm for July 2014 (b). The colour code represents the three different regions identified during the polar dome analysis. Panel (c) shows the distribution of the fraction of particles containing trimethylamine (TMA) with respect to the total amount of particles measured by single particle mass spectrometry (Köllner et al., 2017) in the maximum potential temperature and median latitude coordinate system. Note that for the reason of particle spectra statistics the resolution of median latitude and maximum potential temperature interval is reduced compared to the refractory black carbon distribution (BC) shown in panel (d). The enhancement in BC within the polar dome is most probably due to fresh local pollution.

Having defined the polar dome based on trace gas gradients now allows for a more detailed study on aerosol associated with the different air masses. Efficient wet removal and less efficient transport from lower latitudes lead to generally low aerosol concentrations (Stohl, 2006; Engvall et al., 2008), especially within the Arctic lower troposphere during summer. This is consistent with results in Fig. 15a and b. Observations of elevated levels of accumulation mode particles ($N_{>100}$) can be associated
with regions outside the polar dome and subsequent transport to the measurement region. In parallel, regions within the polar dome were characterized by $N_{\text{>100}}$ smaller than $100 \text{ cm}^{-3}$. In contrary, number concentrations of ultrafine particles ($N_{5-20}$) showed occasionally larger values within the polar dome compared to outside (Fig. 15a), indicating the formation of ultrafine particles occurred within the polar dome region (Burkart et al., 2017). Exemplary for aerosol composition, particulate trimethylamine (measured by single particle mass spectrometry) can be associated with sources within the polar dome (Fig. 15c), consistent with results in Köllner et al. (2017) and Willis et al. (2017). In contrast, the abundance of refractory black carbon can be linked to pollution sources outside the polar dome and subsequent transport to the measurement region (Fig. 15d; Schulz et al., 2019). To conclude, the method introduced in this study is a useful tool to combine Arctic aerosol observations with transport processes and sources within and outside the polar dome region.

Figure 15: The way the aerosol results are presented with the colored boxes makes it difficult to see the shading. Figure 15 is replaced (see new figures above) to demonstrate the applicability of our empirical dome boundary to other constituents or processes using the aerosol formation just as an example.

Technical comments:

Please the comments in the attached PDF manuscript.

Please also note the supplement to this comment:

We thank the reviewer for the technical comments, which were implemented!

Acknowledgement:

All atmospheric data from Zeppelin are publicly available in the EBAS database (http://ebas.nilu.no) and we thank Cathrine Lund Myhre and NILU - Norwegian Institute for Air Research for making the CO and CO$_2$ observations from Zeppelin available.
Characterization of Transport Regimes and the Polar Dome during Arctic Spring and Summer using in-situ Aircraft Measurements

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Abstract. The springtime composition of the Arctic lower troposphere is to a large extent controlled by transport of mid-latitude air masses into the Arctic, whereas during the summer, precipitation and natural sources play the most important role during summer. Within the Arctic region, there exist sloping isentropes creating a transport barrier, known as the polar dome, which results from sloping isentropes. The polar dome, which varies in space and time, and exhibits a strong influence on the transport of air masses from mid-latitudes, enhancing it during winter and inhibiting it during summer. Furthermore, a definition for the location of the polar dome boundary itself is quite sparse in the literature.

We analyzed aircraft based trace gas measurements in the Arctic during two NETCARE airborne field campaigns (July 2014 and April 2015) with the Polar 6 aircraft of Alfred Wegener Institute Helmholtz Center for Polar and Marine Research (AWI), Bremerhaven, Germany. Polar 6 aircraft, covering an area from Spitsbergen to Alaska (134°W to 17°W and 68°N to 83°N). For the spring (April 2015) and summer (July 2014) season we analyzed Using these data we characterized transport regimes of mid-latitude air masses travelling to the high Arctic based on CO and CO₂ measurements as well as kinematic 10-day back trajectories. We found that dynamical isolation of the high Arctic lower troposphere caused by the transport barrier leads to gradients of chemical tracers reflecting different local chemical lifetimes, sources and sinks. Particularly, in particular, gradients of CO and CO₂ allowed for a trace gas based definition of the polar dome boundary for the two measurement periods, which showed pronounced seasonal differences. For both campaigns a transition zone rather than a sharp boundary was derived. For, we derived a transition zone from both campaigns. In July 2014 the polar dome boundary was determined to be at 73.5°N latitude and 299 – 303.5 K potential temperature, respectively. During April 2015 the polar dome boundary was on average located at 66 – 68.5°N and 283.5 – 287.5 K. Tracer-tracer scatter plots and probability density functions confirm different air mass properties inside and outside of the polar dome for the July 2014 and
April 2015 data set. Using the tracer derived polar dome boundaries the analysis of aerosol data indicates secondary aerosol formation events in the clean summertime in both spring and summer.

Further, we explored the processes controlling the recent transport history of air masses within and outside the polar dome. Air masses within the spring-time polar dome mainly experienced diabatic cooling while travelling over cold surfaces. In contrast, air masses in the summertime polar dome were diabatically heated due to insolation. During both seasons air masses outside the polar dome slowly descended into the Arctic lower troposphere from above through radiative cooling. Ascent to the middle and upper troposphere mainly took place outside the Arctic, followed by a northward motion. Air masses inside and outside the polar dome were also distinguished by different chemical composition of both trace gases and aerosol particles. We found that the fraction of amine containing particles, originating from marine biogenic sources, is enhanced inside the polar dome. In contrast, concentrations of refractory black carbon are highest outside the polar dome indicating remote pollution sources.

Synoptic-scale weather systems frequently disturb this transport barrier the transport barrier formed by the polar dome and foster exchange between air masses from mid-latitudes and polar regions. During the second phase of the NETCARE 2014 measurements a pronounced low pressure system south of Resolute Bay brought inflow from southern latitudes that, which pushed the polar dome northward and significantly affected trace gas mixing ratios in the measurement region. Mean CO mixing ratios increased from 77.9 ± 2.5 ppbv to 84.9 ± 4.7 ppbv from the first period to the second period between these two regimes. At the same time CO₂ mixing ratios significantly dropped decreased from 398.16 ± 1.01 ppmv to 393.81 ± 2.25 ppmv.

We further analysed processes controlling the recent transport history of air masses within and outside the polar dome. Air masses within the spring-time polar dome mainly experienced diabatic cooling while travelling over cold surfaces. In contrast, air masses in the summertime polar dome were diabatically heated due to insolation. During both seasons air masses outside the polar dome slowly descended into the Arctic lower troposphere from above caused by radiative cooling. The ascent to the middle and upper troposphere mainly took place outside the Arctic, followed by a northward motion. Our results demonstrate the successful application of utility of applying a tracer based diagnostic to determine the location of the polar dome boundary for interpreting observations of atmospheric composition in the context of transport history.

1 Introduction

In recent decades the Arctic has undergone dramatic changes affecting sea ice, snow, permafrost, surface temperature, land, snow—and atmospheric circulation (IPCC, 2013). Rising temperatures, twice as fast as in the rest of the world, lead have led to a significant retreat of Arctic sea ice (Stroeve et al., 2012; Jeffries et al., 2013). In addition to the reduced extent the thickness of sea ice is continuously decreasing (Lindsay and Schweiger, 2015). The continuing retreat of Arctic sea ice will increase the accessibility of the Arctic thus leading to a potential increase of emissions from local sources of pollutants like shipping (Eckhardt et al., 2013; Corbett et al., 2010; Melia et al., 2016) and oil and gas extraction (Peters et al., 2011). Already to date atmospheric pollutants such as aerosol particles and tropospheric ozone contribute to Arctic warming (Shindell and Faluvegi, 2009). It was shown by earlier studies that mid latitude emissions has continuously decreased (Lindsay and Schweiger, 2015).
Mid-latitude emissions in the Northern Hemisphere are still the main source region for atmospheric pollutants in the Arctic (Barrie, 1986; Koch and Hansen, 2005; Stohl, 2006; Sharma et al., 2013; Arnold et al., 2016). Several studies either based on in-situ measurements or modelling reported enhanced pollution throughout the Arctic troposphere that is dominated by northern Eurasian sources in the lower troposphere and mid-latitude North America and Asia above (Sharma et al., 2006; Shindell et al., 2008; Fisher et al., 2011) even found Asian pollution in the lowermost stratosphere. However, local emissions in specific regions within the Arctic are already important (Stohl et al., 2013) and might gain influence in the near future (Corbett et al., 2010; Peters et al., 2011). Continuing retreat of Arctic sea ice will increase the accessibility of the Arctic, thus leading to a potential increase of emissions from local sources of pollutants like shipping (Eckhardt et al., 2013; Corbett et al., 2010; Melia et al., 2016) and oil and gas extraction (Peters et al., 2011). Resulting atmospheric pollutants, such as aerosol and tropospheric ozone, contribute to Arctic warming (Shindell and Faluvegi, 2009; AMAP, 2015).

Compared to other regions of the Northern Hemisphere, the faster pace of the rising surface and lower tropospheric temperatures in the Arctic is commonly known as Arctic amplification (Holland and Bitz, 2003; Screen and Simmonds, 2010). The interplay between different processes fosters feedback mechanisms that further amplify changes in the environment. The decrease in sea ice reduces the surface albedo and increases latent and sensible heat fluxes into the atmosphere which in turn results in warmer surface temperatures relative to mid-latitudes (Robock, 1983; Hall, 2004; Winton, 2006). Furthermore, Pithan and Mauritsen (2014) reported that temperature feedbacks also play an important role for Arctic amplification. Arctic Amplification could further cause important changes in the mid-latitude circulation (Cohen et al., 2014; Pithan et al., 2018). Zonal winds might weaken and the Rossby wave amplitude is supposed to increase especially during the fall and winter months (Francis and Vavrus, 2012; Francis et al., 2017). Unusual warm sea surface temperatures and low sea ice concentrations in the Arctic have already caused atmospheric circulation anomalies in winter (Lee et al., 2015). Hence, as a result, transport pathways for aerosol and pollution into the Arctic in general will change due to the changing circulation pattern in association with Arctic amplification.

To date it is well known that transport into the Arctic and especially into the Arctic lower troposphere is possible along different pathways, depending on the source area of air masses and the time of the year (Klonecki, 2003; Stohl, 2006; Law and Stohl, 2007; AMAP, 2015). Stohl (2006) identified three major pathways, which significantly contribute to transport from major pollution sources into the Arctic lower troposphere: 1) Rapid low level transport which is followed by an uplift at the Arctic front at the location and time when the Arctic front is located far north. For this transport route uplift and potential precipitation occurs mostly north of 70°N which allows for significant deposition of aerosol and water-soluble pollutants in the Arctic. In their study Stohl (2006) they estimated a transport time of 4 days or less. Significant emissions only from densely populated regions in Europe are able to be transported into the high Arctic lower troposphere via this route since major emission regions in North America and Asia are located south of the polar front. Note that the Arctic front and the polar front are geographically two distinct features. The Arctic front, which is best expressed during the summer months, is thought to develop due to
strong differential heating between the cold Arctic ocean and adjacent ice and snow free land (Serreze et al., 2001; Crawford and Serreze, 2015). The Arctic front marks the southern boundary of the cold Arctic air mass that is separated from the less cold polar air mass at the Arctic front further south. The polar front in contrast is the well known frontal zone separating warm mid-latitude and subtropical air masses from colder polar air masses. It is in general located further south compared to the Arctic front and displaced in equatorward direction in summer and in poleward direction in winter. In this baroclinic region characterized by strong horizontal temperature gradients, cyclones develop from an initial disturbance at the front. During the winter months the Arctic front can extend far south over the continents and can eventually be co-located or merge with the polar front.

2) Low level transport of already cold air masses into the polar dome, which is associated with further diabatic cooling during the transport time scales of 10-15 days. This pathway from European and high latitude Asian sources mainly occurs during winter, since transport over snow-covered regions (e.g. Siberia) is involved. Thus, strongly polluted air masses could be transported into the high Arctic lower troposphere. This transport pathway is negligible during the summer months when the surface in Eurasia is a net source of heat (Klonecki, 2003).

3) Fast uplift mainly due to convection in southern mid-latitudes followed by high altitude transport in northerly directions. Radiative cooling eventually leads to a slow descent into the polar dome area after air masses have arrived in the high Arctic. Being less frequent from Europe, this transport pathway is most prevalent from North America and East Asia. In contrast to the other two transport pathways, scavenging processes can occur during the strong ascent in mid-latitudes which can lead to a significant washout of aerosol and soluble pollutants already outside the Arctic.

The high Arctic lower troposphere is quite well isolated from the rest of the Arctic by a transport barrier referred to as the polar dome. The polar dome is formed by sloping isentropes, the isolines of potential temperature Θ, as a result of radiative cooling in the high Arctic, especially during the winter months without sunlight (Barrie, 1986; Klonecki, 2003; Stohl, 2006). Air masses preferably keep their potential temperatures almost constant during transport, since atmospheric circulation can be well described by adiabatic motions in the absence of diabatic processes related to clouds, radiation and turbulence. The potential temperature is low within the polar dome area and thus only air masses which have experienced diabatic cooling are able to enter the polar dome from specific source regions as discussed before through the pathways discussed above (Stohl, 2006). As a consequence the commonly known “Arctic haze” phenomenon is mainly fed by northern Eurasian pollution sources as those air masses are cold enough to enter the high Arctic lower troposphere (Carlson, 1981; Rahn, 1981; Raatz, 1985; Iversen, 1984; Barrie, 1986; Brock et al., 1990; Dreiling and Friederich, 1997). Already known for decades, Arctic haze has again gained attention at the beginning of the 21st century which was mainly triggered by black carbon (BC) and its role in Arctic climate change (Flanner et al., 2007; Hansen and Nazarenko, 2004; Law and Stohl, 2007; McConnell et al., 2007; Quinn et al., 2008; Shindell and Faluvegi, 2009). Pollution originating from outside the Arctic is transported into the high Arctic lower troposphere during the winter months and the lack of sunlight allows for a build-up of aerosol particles and gaseous pollutants. When temperatures during the winter months become extremely low near the surface, the Arctic lower troposphere is thermally very stably stratified accompanied by surface based inversions that can persist for several days (Bradley et al., 1992). Turbulent exchange and hence dry
deposition is reduced under these conditions. Furthermore, the lower troposphere is extremely dry, preventing scavenging of aerosol and gaseous pollutants by wet deposition. At the spring-time peak, Arctic haze is often visible as layers of brownish haze affecting the radiation budget of the Arctic lower troposphere and also contributing to contamination of the Arctic environment. During the transition to pristine summer conditions Arctic haze declines which is mainly caused by efficient aerosol scavenging in mid-latitudes during convective uplift of air masses. Anthropogenic aerosol is further reduced by frequent precipitation of low intensity within the Arctic lower troposphere (Barrie, 1986; Browse et al., 2012; Garrett et al., 2010).

In general, the polar dome boundary, acting as a transport barrier for warmer mid-latitude air masses, is variable in time and space. Synoptic disturbances can lead to a shift of the polar dome boundary or perturb the transport barrier, fostering and foster exchange with mid-latitude air that can alter the composition of the lower Arctic troposphere. A distinct definition of the polar dome boundary location is crucial to understand and quantify these effects on atmospheric composition. Although the polar dome feature is known for decades, only very few specific definitions of the polar dome boundary have been described in the literature. Early studies by Carlson (1981) or Raatz (1985) identified the polar front as a transport barrier decoupling the Arctic from influence of mid-latitude air masses for their analyses of Arctic haze.

More recent studies used the location of the Arctic front as a marker for the polar dome as a transport barrier (Klonecki, 2003; Stohl, 2006). Jiao and Flanner (2016) used the maximum zonal mean latitudinal gradient of 500 hPa geopotential height in the Northern Hemisphere. One of the drawbacks of the latter approach is the missing definition at lower altitudes.

It was previously mentioned that the polar dome is well isolated from the surrounding troposphere. This leads, which lead to long residence times of air masses within the polar dome. Anthropogenic tracers like CO and CO2 show temporal changes within days to weeks due to changes in emissions and thus, the source strength of these species. Furthermore, the distribution of sources and sinks as well as the efficiency of removal processes for both species is different within the Arctic and at mid-latitudes, leading to latitudinal gradients for both species. Taking into account the isolation created by the polar dome, tracer gradients across the polar dome boundary as a transport barrier should establish measurable gradients in tracer species across its boundary. In this study we use trace gas gradients of CO and CO2 between inside and outside of the polar dome to derive a tracer based diagnostic to identify the location of the polar dome boundary location. The basis for this analysis are two airborne field campaigns: NETCARE 2014 in July and NETCARE 2015 in April (Section 2, 3, 4). Thus, the time period of the late spring and summer are covered, covering late spring to summer. Despite focusing on only these specific time periods, this study is the first attempt to define the polar dome boundary based on airborne trace gas gradients (Section 5). Furthermore, an analysis of the transport history of air masses in We use our trace gas definition of the polar dome boundary to analyze transport history and atmospheric composition within and outside the polar dome and the surrounding is presented.
Figure 1. Compilation of flight tracks of research flights during two NETCARE airborne field campaigns in July 2014 and April 2015.

Table 1. Location and coordinates for the different stations from which measurement flights were performed during the two NETCARE airborne projects in July 2014 and April 2015. Additionally the time at the station and the number of research flights are given.

<table>
<thead>
<tr>
<th>Location</th>
<th>Coordinates</th>
<th>Date</th>
<th>Flights</th>
</tr>
</thead>
<tbody>
<tr>
<td>NETCARE 2014</td>
<td>Resolute Bay</td>
<td>74.7°N, 95.0°W</td>
<td>July 4th to July 21st</td>
</tr>
<tr>
<td>NETCARE 2015</td>
<td>Longyearbyen</td>
<td>78.2°N, 15.5°E</td>
<td>April 5th</td>
</tr>
<tr>
<td></td>
<td>Alert</td>
<td>82.5°N, 62.3°W</td>
<td>April 7th to April 9th</td>
</tr>
<tr>
<td></td>
<td>Eureka</td>
<td>80.0°N, 85.8°W</td>
<td>April 11th to April 17th</td>
</tr>
<tr>
<td></td>
<td>Inuvik</td>
<td>68.3°N, 133.5°W</td>
<td>April 20th to April 21st</td>
</tr>
</tbody>
</table>

2 The NETCARE project

The NETCARE project (Network on Climate and Aerosols: Addressing Key Uncertainties in Remote Canadian Environments, http://www.netcare-project.ca) is configured around four research activities addressing key uncertainties in the field of Arctic aerosol climate research (Abbatt et al., 2019). Within this framework two aircraft based measurement campaigns were performed in the high Arctic. The main objectives of both campaigns were to study aerosol-cloud interactions as well as to characterize local and remote sources for pollution within the high Arctic lower troposphere in summer 2014 (July) and spring 2015 (April). Figure 1 shows a compilation of flight tracks for the two airborne research activities named NETCARE 2014 and NETCARE 2015.

The first project was performed from July 4th to July 21st, 2014 with the Polar 6 aircraft based in Resolute Bay, Nunavut, Canada (e.g. Aliabadi et al. (2016); Leaitch et al. (2016); Willis et al. (2016, 2017); Burkart et al. (2017); Köllner et al. (2017)). In total 11 research flights, each between 4-6 hours long, covered two main research areas, the areas, Lancaster Sound east of
Resolute Bay and the area north of Resolute Bay where two polynyas were located. During the last part of the campaign, July 19th to July 21st, a special research focus was on ship emission measurements (Aliabadi et al., 2016).

The second aircraft project took place in April, 2015. We performed pan-Arctic measurements throughout the European and Canadian Arctic (see Fig. 1). This campaign was a joint NETCARE and PAMARCMiP (https://www.awi.de/en/science/climate-sciences/sea-ice-physics/projects/netcare-arctic-study-of-short-lived-climate-pollutants/pamarcmip-2015.html) project, which will be referred to as “NETCARE 2015” throughout this paper (e.g. Libois et al. (2016); Willis et al. (2019); Schulz et al. (2018) Libois et al)

During 10 research flights, each 4-6 hours long, we specifically focused on a better understanding of aerosol transport into the Arctic in early spring and its influence on ice cloud formation. More details of the different base stations can be found in Tab. 1. Multiple vertical profiles from the lowest possible altitude (of 60 m) up to 6000 m were performed to study the vertical distribution of aerosol particles and trace gases.

3 Methodology

3.1 Measurements and data platform

Airborne measurements were performed using the Polar 6 aircraft of the Alfred Wegener Institute Helmholtz Center for Polar and Marine Research, Bremerhaven, Germany. Polar 6 is a DC 3 aircraft converted to a Basler BT67 (Herber et al., 2008) and modified for operation in cold and harsh environments. The aircraft has a non-pressurized cabin, however flights up to an altitude of 6 km were regularly performed during the 2014 and 2015 campaigns. The typical survey speed of the aircraft is 120 kts (∼60 m s⁻¹) with ascent and descent rates of 150 – 300 m min⁻¹ during the vertical profiles.

3.2 Instrumentation

Meteorological and aircraft altitude data for Polar 6 are provided by the AIMMS-20 instrument. The instrument was designed and manufactured by Aventech Research Inc., Barrie, Ontario, Canada. It includes the Air Data Probe (ADP) that reports the three-dimensional, aircraft-relative flow vector consisting of true air speed, angle-of-attack and sideslip. In the rear section of the instrument, temperature and relative humidity sensors are located providing data with an accuracy of 0.30°C and a resolution of 0.01°C for temperature measurements and 2.0 % and 0.1 % for humidity measurements, respectively. A GPS module provided the aircraft 3-D position and inertial velocity. Horizontal and vertical wind speeds were measured with accuracies of 0.50 and 0.75 m s⁻¹, respectively. All data were internally sampled with 200 Hz resolution and for further analysis averaged to 1 Hz resolution. From the AIMMS-20 data set, especially the temperature and pressure data are used throughout this study. The instrumentation for aerosol and cloud droplets as well as upwelling radiance measurements is described in detail in Leaitch et al. (2016); Willis et al. (2016); Burkart et al. (2017); Aliabadi et al. (2016); Libois et al. (2016) and Schulz et al. (2018) Schulz et al. (2019).

CO was measured with an Aerolaser ultra fast carbon monoxide (CO) monitor model AL 5002 based on VUV-fluorimetry, using the excitation of CO at 150 nm (Gerbig et al., 1999). UV light stems from a resonance lamp excited by a Radio Frequency
(RF) discharge. The selection of the 150 wavelength is realised by an optical filter, which images the lamp into the RF chamber, where fluorescence is viewed at a right angle by means of a photomultiplier tube (PMT) with suprasil optics. The optical filter consists of two lenses. The two dielectric mirrors provide the spectral band path (bandwidth of 8 full width at half maximum (FWHM) at approximately 150). The Details on the measurement principle can be found in Gerbig et al. (1999) or Scharffe et al. (2012). The instrument was modified for applying in-situ calibrations during in-flight operations. These We performed these regular in-situ calibrations are performed on a 15 to 30 min time interval during measurement flights using a NIST traceable calibration gas with a known CO concentration at atmospheric levels as well as zero measurements. Calibrations and zero measurements account for instrument drifts. CO data achieved a precision (1σ, 1 Hz) of 2.2 ppbv during NETCARE 2014 and 1.5 ppbv during NETCARE 2015. The stability of the instrument is calculated to 4.1 ppbv and 1.7 ppbv, respectively, before applying the post flight data correction. Stability is a measure of reproducibility and based on the mean drift between two subsequent calibrations which were performed during flights. Stability is mainly affected by temperature variations. These instrumental drifts are corrected after the flights assuming linear drift. Hence, the total uncertainty of 4.7 ppbv relative to the working standard of 4.7 for NETCARE 2014 and 2.3 ppbv for NETCARE 2015 can be regarded as an upper limit.

CO₂ was measured with a LI-7200 closed CO₂/H₂O Analyzer from LI-COR Biosciences GmbH. The instrument simultaneously also measures water vapour, which is used for CO₂-H₂O-interference corrections. The measurement principle is based on an optical source emitting infrared light through a chopper filter wheel and the enclosed sample path to a temperature controlled lead selenide detector. By using the ratio of absorption by carbon dioxide in the sample path to a reference, the density of the gases and thus the mixing ratio can be calculated. The instrument itself was mounted in a 49â°19″, 3 HE rack mount including additional components for flow control and in-situ calibrations during in-flight operations. As for the CO measurements, calibrations were performed we performed calibrations on a regular time interval of 15 to 30 minutes using a NIST traceable calibration gas with a known CO₂ concentration at atmospheric levels and a water vapour concentration close to zero. CO₂ data during NETCARE 2014 achieved a precision (1σ, 1 Hz) of 0.02 ppmv and 0.05 ppmv during NETCARE 2015. Using the same methodology as for CO, the stability of the instrument is calculated to 0.76 ppmv for NETCARE 2014 and 1.72 ppmv for NETCARE 2015, before applying the post flight data correction. The total uncertainty relative to the working standard thus amounts to 0.76 ppmv for NETCARE 2014 and 1.72 ppmv for NETCARE 2015. The uncertainty for the measurement of H₂O is 18.5 ppmv or 2.5 %, whichever is greater.

3.3 LAGRANTO backward trajectories

We used the Lagrangian analysis tool (LAGRANTO) (Wernli and Davies, 1997; Sprenger and Wernli, 2015) to determine the origin of air masses that were sampled. LAGRANTO trajectories were calculated based on operational analysis data from the European Centre of Medium-Range Weather Forecasts (ECMWF). This data has a horizontal grid spacing of 0.5° with 137 hybrid sigma-pressure levels in the vertical from the surface up to 0.01 hPa. Trajectories were initialized every 10 s from coordinates along individual research flights and calculated 10 days back in time. The location of the individual trajectory is available at a 1 h time interval. Different variables of atmospheric state were simulated along the trajectory (temperature,
potential temperature, potential vorticity, specific humidity, cloud water and cloud ice water content, Richardson number and equivalent potential temperature).

To account for the latitudinal transport history of the air parcels we calculated the median latitude along the trajectories. We used this as a proxy for the most representative position during the last 10 days associated with the respective values of CO and CO₂. To account for diabatic descent occurring during transport we calculated the maximum potential temperature. Both parameters are used in Sec. 5.2 for the analysis of the polar dome boundary. As a measure for the uncertainty of the temperature along the trajectory we calculated the median difference between temperatures measured in-situ on the aircraft and the corresponding temperatures interpolated to the initialisation point of the trajectory along the flight track based on analysis data. For the measurements in July 2014 the median difference is 0.31 °C (interquartile range: −0.71 – 1.72 °C). For the April 2015 measurements, the respective median difference is 1.50 °C (interquartile range: 0.69 – 2.14 °C).

4 Meteorological Overview

4.1 NETCARE 2014

Flights during the NETCARE 2014 field campaign were performed during Arctic summer in July in the area around Resolute Bay, Nunavut, Canada. The meteorological situation can be separated into two different meteorological regimes (see Fig. 2). During the first phase (July 4th to July 12th) the boundary layer was capped at low altitudes by a distinct temperature inversion leading to a very stable stratification of the lower troposphere. The prevailing influence of a high pressure system provided ideal conditions for aircraft based measurements with mainly clear sky, only few or scattered clouds and low wind speed. Beginning July 13th Resolute Bay was influenced by a low pressure system located to the west above the Beaufort Sea. This system eventually passed Resolute Bay two days later. Increased humidity, precipitation and fog characterized the local weather and prevented Polar 6 from flying until July 17th. The last flights of the campaign were performed between July 19th and July 21st when a pronounced low pressure system south of Resolute Bay and centred around King William Island influenced the measurement region (see Fig. 2b). Increased wind speeds, mostly mid to high level clouds and precipitation resulted from the inflow of warm air from more southern latitudes. Furthermore, this situation was favourable for mid-latitude air masses being advected to the measurement region potentially affecting concentrations levels of trace gases and aerosol particles.

4.2 NETCARE 2015

The NETCARE 2015 pan-Arctic study was performed during Arctic spring in April in the European and Canadian Arctic. The aircraft were based at four different locations, namely Longyearbyen (Norway), Alert, Eureka (both Nunavut, Canada) and Inuvik (Northwest Territories, Canada), allowing for a wider coverage of the entire Arctic. Figure 3 shows mean geopotential height at 850 hPa over the time interval of the measurements in the respective region. At the time of the first flight of the campaign in Longyearbyen (April 5th, see Fig. 3a), Spitsbergen was under a quite stable high pressure influence with almost no clouds and only weak winds. During the measurements in Alert the meteorological situation was dominated by a pool
of cold air centred above Ellesmere Island to the south-west of Alert. A cyclonic flow was established around this cold air guiding low pressure systems around the cold pool and thus preventing mid-latitude air masses to potentially influence the high Arctic lower troposphere. Stable conditions with almost clear sky facilitated airborne measurements on 4 research flights between April 7\textsuperscript{th} and April 9\textsuperscript{th} during this period (see Fig. 3b). After the transfer to Eureka on April 10\textsuperscript{th} two research flights were performed in almost the same meteorological conditions as in Alert. When the surface low started moving south over Baffin Bay from April 13\textsuperscript{th} onward (see Fig. 3c) strong northerly and north-easterly winds in the lower troposphere influenced the measurement regions. The warmer flow was guided from southern areas over open water around the low pressure centre and was associated with moisture transport to the land leading to cloud formation and fog, which impeded research flights out of Eureka. In the following days a low pressure system started intensifying north of Greenland and maintained the low level moist northerly flow. The last flights of the campaign were conducted in Inuvik between April 20\textsuperscript{th} and April 21\textsuperscript{st}. After the ferry from Eureka to Inuvik on April 17\textsuperscript{th} and 18\textsuperscript{th} high pressure influence was prevalent in Inuvik. At the time of the research flights a low pressure system located over Alaska fostered a southerly and south easterly flow into the Inuvik area, favourable for mid-latitude air masses to enter the measurement region (see Fig. 3d). A detailed description of the air mass history corresponding to the meteorological situation is given in the supplementary material (Fig. S1 and S2).

5 Results

5.1 Air-mass-history

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**Figure 2.** Mean geopotential height on 850 hPa in geopotential decameter (gpdm) for the period from July 4\textsuperscript{th} 2014 to July 12\textsuperscript{th} 2014 (a) and for the period from July 17\textsuperscript{th} 2014 to July 21\textsuperscript{st} 2014 (b).
Figure 3. Mean geopotential height on 850 hPa in geopotential decameter (gpdm) from April 5th 2015 (a), from April 7th 2015 to April 9th 2015 (b), from April 11th 2015 to April 13th 2015 (c) and from April 20th 2015 to April 21st 2015 (d). Our flight location for each time period is marked in yellow.

Area weighted trajectory density during the different campaign phases in July 2014 (gridded by 1° x 1°). The colour code represents the amount of trajectory points per grid box weighted by the area of the grid box. The individual panels show the results for the first phase (a) and the second phase (b). The bold black dashed circle denotes the Arctic circle.

During the flights of the first period of NETCARE 2014 until July 12th high pressure influence was prevailing. Air masses tended to stay within the high Arctic and circle around the measurement region. Almost no mid-latitude influence in terms of trajectory origin was observed when using the Arctic circle as a boundary between the Arctic and mid-latitudes. This is evident...
in Fig. 4a, which shows the area-weighted accumulated number of trajectory points per grid box (1° x 1°) for the last ten days before the measurements during the summer campaign in July 2014. The highest density of trajectory points is observed significantly north of the Arctic circle. Note that multiple "hits" of one trajectory in a specific grid box are possible during the 10-day travel of the air mass associated with the trajectory.

Area-weighted trajectory density during the different campaign locations in April 2015 (gridded by 1° x 1°). The colour code represents the amount of trajectory points per grid box weighted by the area of the grid box. The individual panels show the results for the measurements in Longyearbyen (a), Alert (b), Eureka (c) and Inuvik (d). The bold black dashed circle denotes the Arctic circle.

In contrast, during the flights within the second period (July 17th to July 21st) more air masses originated in regions south of the Arctic circle (see Fig. 4b). Highest trajectory densities are found slightly north of the Arctic circle in the Canadian Arctic Archipelago extending southward to continental Canada and the Bering Sea. The stable low-pressure system over King William Island thus favours the transport of mid-latitude air masses to the high Arctic, and the potential for a stronger impact of mid-latitude sources on the Arctic chemical composition.

During pan-Arctic measurements in April 2015, air masses show very different histories depending on the respective measurement location in the Arctic (see Figs. 5a-d). Measurements performed from the two northernmost stations Alert and Eureka are less influenced by air masses of mid-latitude origin than those further south. This indicates that synoptic disturbances did not have a strong influence on the high Arctic stations at least during the one-month period, which is covered by our measurements and the backward trajectories. Based on the "density maps" of the trajectory locations, the strongest mid-latitude influence is indicated during the measurements in Inuvik between April 20th and April 21st (Fig. 5d).

During three flights, a warm conveyor belt (WCB) type transport associated with a strong low-pressure system over Alaska influenced the tropospheric composition in the measurement region by advecting pollution from South East Asia. As already mentioned before the high Arctic stations in Eureka and Alert were much less affected by air masses from mid-latitudes, which is confirmed by the analysis of the air mass history for the flights between April 7th and April 13th (Figs. 5b and c) when only a few trajectories travel over areas outside the Arctic circle. Air masses mainly resided over the Canadian Arctic Archipelago and northern Greenland with only episodic influence from the North American continent and Siberia. In Longyearbyen only one flight was performed on April 5th, which shows a mixture of mid-latitude and Arctic air masses (Fig. 5a). The origin of the majority of air masses contributing to the observations in Longyearbyen was in northern and eastern parts of Europe.

5.1 Trace gas observations

According to the meteorological situation and the general transport regimes discussed above, we expect a significant influence of the air mass history on the mixing ratios of CO and CO2. Both species show latitudinal and vertical gradients (see Fig. 6) and in addition, one is affected by anthropogenic pollution which makes them ideally suited for identification of pollution events affecting the Arctic background. Furthermore, both species show a seasonal cycle in the Arctic.
Figure 4. CO (Dlugokencky et al., 2018) (a) and CO₂ (Petron et al., 2018) (b) seasonal cycle based on NOAA ground based measurements in Alert (Canada), Barrow (Alaska) and Mace Head (Ireland) for the years 2014 and 2015. Running means are shown for the respective station data (symbols). Mean aircraft data for altitudes lower than 200 m for individual flights are overlaid. Error bars (yellow and orange shading) for the aircraft data are too small to be visible. NETCARE 2014 data are in yellow and NETCARE 2015 data are in orange.

(a) and (b) seasonal cycle based on NOAA ground based measurements in Alert (Canada), Barrow (Alaska) and Mace Head (Ireland) for the years 2014 and 2015. The running mean line colours correspond to those of the station symbols. The aircraft data of the lowest 200 are overlayed as mean plus standard deviation for the respective flight. Error bars for the aircraft data are too small to be visible. NETCARE 2014 data are in yellow and NETCARE 2015 data are in orange.

Figure 6 shows the ground based observations of (Dlugokencky et al., 2018) (a) and (Petron et al., 2018) (b) for three relevant sites, namely Alert (Canada, 82.4 °N), Barrow (Alaska, 71.3 °N) and Mace Head (Ireland, 53.3 °N) for the years 2014 and 2015. Superimposed are the respective trace gas Our aircraft based measurements of CO and CO₂ for altitudes below 200 as mean values for each individual flight during the two NETCARE campaigns in July 2014 and April 2015. For and aircraft based and ground based observations show are in very good agreement with ground based observations (see Fig. 4). In April 2015 trace gas levels are in general higher compared to July 2014. The observed change in trace gas levels between spring and summer reflects the typical seasonal cycle of these two species in the Arctic.

For the high Arctic CO the seasonal cycle in the high Arctic seasonal cycle shows a maximum in late winter/early spring and a minimum during late summer. This seasonal cycle maximum reflects quite well the transport of anthropogenic pollutants - mainly from fossil fuel burning - from northern Europe and Siberia into the high Arctic lower troposphere during winter (Klonecki, 2003; Stohl, 2006). Since photochemically produced OH is absent during wintertime due to the lack of sunlight, CO in the high Arctic has no significant sink which results in a longer chemical lifetime of the order of months. Hence, CO increases over the course of the winter, in particular within the polar dome (Novelli et al., 1998; Engvall et al., 2008). As soon as sunlight returns during late February and early March there is a sharp transition...
between 24 h polar night and 24 h polar day. The increasing concentration of OH leads to increased oxidation of CO and a shorter lifetime in the order of weeks (Dianov-Klokov and Yurganov, 1989; Holloway et al., 2000). During the transition from spring to summer (April to June) the photochemical activity in the Arctic and smaller mid-latitude emissions of CO in mid-latitudes lead to decreasing CO in the Arctic until a minimum is reached at the end of the summer (Barrie, 1986; Klonecki, 2003; Engvall et al., 2008).

The seasonal cycle for CO₂ in the northern hemisphere is mainly controlled by carbon uptake and release processes of the biosphere (Keeling et al., 1996; Forkel et al., 2016). Whereas during the summer months the CO₂ seasonal cycle reaches its minimum due to photosynthetic carbon uptake by vegetation, respiration of the biosphere is prevalent during wintertime. Particularly in Arctic winter, the absence of sunlight allows for a build-up of CO₂ concentrations. However, meridional CO₂ transport into the high Arctic by synoptic weather disturbances plays a critical role for the seasonal cycle and dominates over local atmosphere-biosphere fluxes (Fung et al., 1983; Parazoo et al., 2011; Barnes et al., 2016). As a result, the synoptic eddy driven meridional transport reduces the seasonal cycle in mid-latitudes and amplifies it in polar regions, leading to a meridional CO₂ gradient (Parazoo et al., 2011).

5.2 The location of the polar dome

Conceptually, the polar dome can be regarded as the region below the upward sloping isentropes north of the region where low-value isentropes intersect, and horizontally as the region north of where low-valued isentropes intersect with the surface. Figure 7-5 shows the observed potential temperature (Θ) distribution as a zonal mean for the respective campaigns. Potential temperature was calculated from temperature and pressure measurements on board the Polar 6 aircraft. A dome-like structure of the isentropes is visible for April 2015 (NETCARE 2015). Minimum potential temperatures lower than 275 K were only present in the high Arctic lower troposphere north of 70°N. In contrast, a dome-like
structure is hardly visible for July 2014 (NETCARE 2014). Only below 950 hPa and north of 75°N Θ values of 275 K were observed. This is in agreement with previous studies which showed that during the summer months the extent of the polar dome is much smaller compared to the winter time (Klonecki, 2003; Stohl, 2006; Jiao and Flanner, 2016).

If we now use potential temperature as the vertical coordinate, air masses within the polar dome associated with the coldest potential temperatures should separate from other regions. This is evident on Fig. 8, in particular separation is evident in Fig. 6, particularly for the April 2015 measurements (b). The colour code in Figs. 8a and b represents the average CO background mixing ratio calculated from all data points within the respective bin interval. For our analysis of the polar dome we only use background trace gas mixing ratios. We exclude polluted air masses, if the mean background CO mixing ratio is exceeded by 2 standard deviations. In April 2015 (Fig. 8b) northernmost latitudes exhibited the largest CO values of 140-150 ppbv for potential temperatures lower than 275 K. At higher isentropes typical CO values range from 100-135 ppbv. There, at these higher potential temperatures, the larger variability of CO mixing ratios indicates different source regions contributing to the observations, which is in particular observed for the lower latitudes. In the distribution for the summer campaign in July 2014 (Fig. 8a) a region of rather uniform low CO is evident north of 75°N and below 290 K. Mixing ratios south of 75°N and above 290 K tended to be more variable and in general larger than within the aforementioned northern region. Again, the increased variability results from different air mass origins associated with different levels of CO. For both measurement campaigns, we observed a distinct transition between the northernmost lower troposphere and regions with lower latitudes and larger potential temperatures is observed. This transition indicates a transport barrier for air masses to reach the high Arctic lower troposphere. We hypothesize that those regions north of 75°N showing the lowest CO mixing ratios during July 2014, and the largest during April 2015, represent the polar dome whereas the rest of the measurements were collected outside the polar dome.

**Figure 6.** CO distribution binned by latitude and potential temperature for July 2014 (a) and April 2015 (b). The colour code represents the average CO mixing ratio calculated from all data points in the respective 1° latitude and 2 K bin interval. Note that only background mixing ratios are shown now. Polluted air masses are identified and filtered when the average background distribution is exceeded by 2σ.
Figure 7. (a-d): Trace gas distribution in the maximum potential temperature and median latitude coordinate system. The median and maximum values were derived from every 10-day-trajectory calculated along the flight track. The colour code is representative for the average CO (CO₂) value calculated from all data points within one bin interval. The trace gas mixing ratio is the measured value which is assumed to stay constant along the respective trajectory for each measured data point every 10 s.

The trace gas distribution in Fig. 8–6 only shows a snapshot of the actual situation at the time and the location of the measurement. We used ten day backward trajectories to take into account different transport pathways to the Arctic and the residence times of air masses inside the polar dome area, which is expected to be potentially higher in that region than residence times in areas outside the polar dome. We display the CO and CO₂ distribution in a maximum potential temperature versus median latitude coordinate system. Maximum potential temperature and median latitude were derived along every individual trajectory. Median latitude allows for a separation between high Arctic air masses and air masses from mid-latitudes. Air parcels isolated in the polar dome region should stay at high median latitudes whereas air masses extending over a larger meridional distance thus show a more southern median latitude. Maximum potential temperature further allows us to account for diabatic descent that air masses experienced during transport. In the maximum potential temperature vs. median latitude coordinate system those air masses inside the polar dome region exhibit the lowest maximum
potential temperatures and at the same time largest median latitudes, and thus separate from air masses outside the polar dome. Furthermore, the polar dome is dynamically well isolated from the surrounding troposphere as discussed earlier. Hence, air parcels inside the polar dome are in general not affected by strong mid-latitude CO sources and should show a relatively small CO variability. Therefore, we remapped the CO data to the median latitude and maximum potential temperature along the trajectory to identify transport regimes and the effect of the transport barrier at the polar dome (see Figs. 97a and c). The majority of trajectories with relatively low CO mixing ratios are confined by the 305 K isentrope and 70°N as evident in Fig. 97a for July 2014. For April 2015 (Fig. 97c) an area with relatively higher CO is located north of a latitude of 65°N and below a potential temperature of around 280 K. These findings are supported by the CO₂ distribution in the same coordinate system displayed in Figs. 97b and d, which show similar boundaries. Hence, gradients of CO and CO₂ establish at these boundaries.

In this study these chemical gradients are in particular used to derive a tracer based definition of the polar dome boundary for the two NETCARE measurement campaigns during July 2014 and April 2015. The determination of Next, we determine the location of the polar dome boundary based on trace gas gradients is discussed in the following section.

Trace gas gradients

In July 2014 both species–CO and CO₂ show a latitudinal gradient across the isentropes for maximum potential temperature levels of Θ < 305 K (see Fig. 97a and b). In particular, CO₂ shows a strong increase from values around 393 ppmv to 398 ppmv towards high latitudes in the median latitude range of 70°N to 75°N. For CO, a decrease from about 83 ppbv to 78 ppbv in this latitude range is also evident. In contrast to CO₂ the large variability of CO at lower median latitudes reflects a larger variability of potential source regions, which in turn partly masks the CO gradient. Above 305 K trace gas gradients are weak or absent, indicating rapid isentropic mixing from lower latitudes. We now calculated isentropic trace gas gradients in layers of 2 K for the maximum potential temperature as the vertical coordinate in Fig. 97 to derive the horizontal polar dome boundary. Hence, for every 2 K altitude interval we determine the latitude of the strongest trace gas gradient. Below 305 K isentropic trace gas gradients maximize around 73°N. We finally used the median of these maximum gradient latitudes to define the polar dome boundary. If we derive a different median value for the maximum gradient for each of the two species and tracer species, we consider this difference as the range of the polar dome boundary which can be interpreted as a transition zone rather than a sharp boundary between inside and outside of the polar dome. For July 2014 the average horizontal polar dome boundary is a sharp transition at 73.5°N (blue bar in Fig. 408b). The interquartile range denotes to 72.5°N - 77°N.

The strongest vertical gradients of CO and CO₂ were determined at maximum potential temperature values of 299 - 303.5 K (blue bar in Fig. 408a; interquartile range: 297 to 304.5 K). These values for the upper polar dome boundary are relatively high given a surface value of potential temperature of typically 280 K in summer. A close inspection of the CO₂ distribution north of 73.5°N (Fig. 97b) reveals two layers in the high Arctic separated by approximately 285 K. The vertical profile of CO₂ clearly shows the two layers (Fig. 408a). The fact that the vertical profile of CO does not show a clear separation, indicates that rather pristine air masses dominate both layers, which have not experienced strong pollution impact, but rather biogenic impact mainly affecting CO₂. If we additionally use this information, we can separate three distinct
Figure 8. Medians of CO and CO$_2$ as a function of median trajectory latitude and maximum trajectory potential temperature (details see text). Median values for the vertical profiles were only calculated north of 75$^\circ$N for July 2014 (a) and north of 65$^\circ$N for April 2015 (c) because latitudinal gradients indicate a dome boundary north of these median latitudes. At lower latitudes transport and mixing homogenize these gradients (see Fig. 9). For both NETCARE campaigns the median horizontal values were derived only below 300 K (b and d). The blue bar marks the latitude and the potential temperature interval of the strongest change in the tracer mixing ratio, which is interpreted as the transition zone of the polar dome boundary. The shaded area in all figures represents the 1 $\sigma$ standard deviation.

air masses. The region with lowest potential temperatures ($\Theta < 285$ K) has small (large) mixing ratios of CO (CO$_2$) and is mostly isolated from mid-latitude influence. These air masses are most likely remnants of the spring time polar dome and we refer to this as the aged polar dome. Between 285 K and 299 K the air masses still show signatures of the polar dome while also the influence from mid-latitudes increases, indicated by lower CO$_2$ mixing ratios. This region is capped in the vertical by the polar dome boundary spanning from 299 to and 303.5 K. Above and thus outside the polar dome, mixing ratios
Figure 9. (a): Tracer-tracer scatter plot of all data points (background + pollution plumes) within the aged polar dome (blue), the mixing region (green) and outside the polar dome (red) for July 2014. (b): Tracer-tracer scatter plot of all data points (background + pollution plumes) within (blue) and outside (red) the polar dome for April 2015. To separate the different regions the tracer derived polar dome boundaries are used. Boundary values for each region are summarized in Table 2.

Table 2. Maximum potential temperature and median latitude values for the polar dome boundary. Included are also the boundary values used for separating the different regions identified for further analysis.

<table>
<thead>
<tr>
<th>Region</th>
<th>Maximum potential temperature</th>
<th>Median latitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>July 2014, polar dome boundary</td>
<td>299.0 - 303.5 K</td>
<td>73.5°N</td>
</tr>
<tr>
<td>April 2015, polar dome boundary</td>
<td>283.5 and 287.5 K</td>
<td>66.0°N - 68.5°N</td>
</tr>
<tr>
<td>July 2014, aged polar dome</td>
<td>$\Theta_{\text{max}} &lt; 285.0$ K</td>
<td>$\text{Lat}_{\text{med}} &gt; 73.5°$N</td>
</tr>
<tr>
<td>July 2014, mixing region</td>
<td>$285.0 \text{K} \leq \Theta_{\text{max}} &lt; 299.0$ K</td>
<td>$\text{Lat}_{\text{med}} &gt; 73.5°$N</td>
</tr>
<tr>
<td>July 2014, outside polar dome</td>
<td>$\Theta_{\text{max}} &gt; 303.5$ K</td>
<td>$\text{Lat}_{\text{med}} &lt; 73.5°$N</td>
</tr>
<tr>
<td>April 2015, polar dome</td>
<td>$\Theta_{\text{max}} &lt; 283.5$ K</td>
<td>$\text{Lat}_{\text{med}} &gt; 68.5°$N</td>
</tr>
<tr>
<td>April 2015, outside polar dome</td>
<td>$\Theta_{\text{max}} &gt; 287.5$ K</td>
<td>$\text{Lat}_{\text{med}} &lt; 66.0°$N</td>
</tr>
</tbody>
</table>

of both species clearly show characteristics of mid-latitude influence. Similar values to those observed outside the polar dome were also found in the mid-latitude lower troposphere for example at the Mace Head observatory in Ireland (see Fig. 6). A summary of the values for the polar dome boundary and the boundaries of the three different regions for July 2014 can be found in Tab. 2.

The threefold structure of the high Arctic lower troposphere, based on the derived boundary values for each region summarized in Tab. 2, is further evident in the CO-CO$_2$ tracer-tracer correlation in Fig. 9. More precisely, the aged polar dome
(blue dots) seems to be a subset of the mixing region (green dots) indicated by a narrow group of data points at end of the highest CO\textsubscript{2} and lowest CO mixing ratios. The aged polar dome region is furthermore clearly separated from the region outside (red dots). The green dots indicate the influence of mixing between dome air and extra-dome air and correspond to the mixing region in the high Arctic between 285 and 299 K (compare Fig. 9).

The tracer derived polar dome boundary for the April 2015 measurements was on average determined between 66.0°N and 68.5°N (blue bar in Fig. 10d; interquartile range: 65.0°N - 69.5°N) for the latitudinal value and between potential temperatures of 283.5 and 287.5 K (blue bar in Fig. 10c; interquartile range: 280.5 and 291.5 K). Values for the polar dome boundary are also summarized in Tab. 2 for April 2015. During spring the CO-CO\textsubscript{2} tracer-tracer correlation in Fig. 11b indicates at least three distinct branches. The separation of air masses between inside the polar dome and outside is based on the tracer derived polar dome boundary (see Tab. 2). The red branch with the highest CO and CO\textsubscript{2} mixing ratios can be associated to pollution events observed during flights in Inuvik. In contrast, the red branch with highest CO\textsubscript{2} but relatively low CO values corresponds to observations in the unpolluted lower troposphere in the Inuvik region. Both branches are clearly associated to air masses outside the polar dome, since measurements around Inuvik were mostly performed outside the determined polar dome boundary. In contrast the blue branch represents the measurements inside the polar dome.

These data points show different slopes indicating different air mass properties. Within the polar dome region we observe a mixture of air masses which is evident by the relatively broad range of CO\textsubscript{2} and CO values forming a mixing line in Fig. 11b. The lowest CO and CO\textsubscript{2} values inside the polar dome (blue) can be associated to the lowest maximum potential temperatures and thus the highest residence time within the dome area. Air masses with highest CO and CO\textsubscript{2} mixing ratios but still inside the polar dome (blue) originate at lower latitudes. In fact, ground stations in the potential source region two weeks before the time of the measurement campaign show enhanced CO and CO\textsubscript{2} values in the range of the upper branch of the scatter plot of those data points inside the polar dome. The observed mixture of air masses is also reported by Willis et al. (2019) who observed an altitude dependent composition and degree of processing of aerosol in the spring time polar dome. In their study, FLEXPART simulations suggest more southern source regions for those air masses with the highest potential temperatures within the polar dome. Furthermore, Schulz et al. (2018) Schulz et al. (2019) determined an increase of refractive black carbon (rBC) and a decrease of the rBC mass-mean diameter with potential temperature inside the spring time polar dome, which was also associated to different source regions contributing to the observations.

5.3 Air mass statistics and tropospheric composition

In order to analyse the processes dominating the recent transport history of observed air masses, we apply the phase-space diagram introduced by Binder et al. (2017). In this analysis, we determine the maximum change in potential temperature (ΔΘ) as the difference between the potential temperature at the time of the measurement (Θ\textsubscript{0}) and the previous potential temperature minimum or maximum (Θ\textsubscript{min}, Θ\textsubscript{max}) along the trajectory. Depending on which absolute difference is larger, the air mass has either experienced diabatic heating (ΔΘ = (Θ\textsubscript{0} - Θ\textsubscript{min}) > 0) or cooling (ΔΘ = (Θ\textsubscript{0} - Θ\textsubscript{max}) < 0). An analogous analysis is made for the absolute temperature to determine if an air parcel predominantly gained or lost temperature before the measurement. This analysis allows us to cluster the data into four categories shown in Figs. 10a-d and 11a-c. The changes in potential
temperature and temperature along the trajectories, indicated by the clusters, can be associated with processes affecting the respective air mass. Sector 1 \((\Delta \Theta < 0, \Delta T < 0)\) mainly contains air masses that experienced diabatic cooling, which indicates either thermal radiation, evaporation or low level transport over snow or ice covered regions (i.e. cold surfaces). In sector 2 \((\Delta \Theta > 0, \Delta T < 0)\) air masses gained potential temperature, which indicates an ascending air mass that is diabatically heated by, for example, solar radiation or condensation processes. Sector 3 \((\Delta \Theta > 0, \Delta T > 0)\) includes those air masses that experienced both an increase in temperature and potential temperature probably due to solar insolation. Finally, sector 4 \((\Delta \Theta < 0, \Delta T > 0)\) combines air masses that lost potential temperature and gained temperature during transport. These air masses are diabatically cooled and thus experienced a descent. We apply this clustering approach to identify differences between observations in the different regions (polar dome, outside polar dome, etc.) and to study whether we can connect trace gas mixing ratios to the dominant process in a specific region. The different regions were separated based on the tracer derived polar dome boundary (see Tab. 2).

For July 2014, three regions are of particular interest, (1) the aged polar dome, (2) the mixing region and (3) the region outside the polar dome (see Figs. 10a-d). Within the aged polar dome sector 3 dominates, thus solar insolation is important for heating the lowest levels. Air masses residing at the lowest altitude experienced diabatic heating potentially resulting in a slow and shallow convective lift of the air masses prior to the time of the measurement. Equal contributions come from sectors 1 and 4, dominated by diabatic cooling either through descent, low level transport over cold surfaces or evaporation. Almost none of the air masses are in sector 2, thus a significant ascent of air masses within the aged polar dome hardly occurs. In contrast, outside the polar dome area sector 4 dominates and generally diabatic cooling occurs (sector 1 and 4). Thus within the polar dome local near-surface diabatic processes seem to mostly affect the air masses; this effect is also evident in air mass composition. Observed CO\(_2\) mixing ratios are highest in the aged polar dome associated with aged Arctic air and negligible mid-latitude influence. Outside the polar dome, air masses have been transported into the Arctic and descended due to radiative cooling. Associated CO\(_2\) mixing ratios of these air masses are significantly lower and can be attributed to more mid-latitude regions. Air masses associated with sector 4 experience descent once they have reached the high Arctic at higher altitudes. This can be regarded as a typical transport pathway during the summer with a fast uplift of air masses at mid-latitudes within convective and frontal systems followed by a northward movement and finally a descent into the high Arctic lower troposphere. Air masses in the mixing region are even more dominated by sector 4. Within this mixing region, trace gas concentrations still show dome-like characteristics and low level processes seem to dominate over episodes of mid-latitude transport associated with air masses with lower CO\(_2\) concentrations. Results for CO confirm the derived transport history for July 2014. A more detailed analysis of the physical processes along the air mass trajectories can be found in the supplement material (Figs. S3, S4 and S5) to this paper.

In April 2015 the picture is quite different (see Figs. 11a-c). Inside the polar dome diabatic cooling dominates, in particular sector 1. This is caused by diabatic descent due to radiative cooling in the absence of sunlight. In addition, low level transport over cold surfaces significantly contributes to transport into the high Arctic. Associated mixing ratios of CO show rather large values that can be explained by the accumulation of anthropogenic pollution from inner Arctic and high northern mid-latitude sources during the winter months. In the absence of sunlight chemical loss is reduced leading to an increase
Figure 10. Phase-space diagram illustrating the maximum absolute change in temperature ($\Delta T$) and potential temperature ($\Delta \Theta$) relative to the time of the measurement for July 2014. The colour code denotes the CO$_2$ mixing ratio at the time of the measurement. (a) shows all background data, (b) shows only those data corresponding to the aged polar dome, (c) shows the data points within the mixing region whereas (d) includes all data points outside the polar dome. To separate the different regions the tracer derived polar dome boundaries are used (see Tab. 2).

in the CO atmospheric lifetime. Air masses within the polar dome are quite efficiently isolated from any significant southern mid-latitude influence. In contrast, outside the polar dome the picture is more diverse. Sector 4 dominates, thus diabatically cooled air masses potentially associated with the tendency to descent are observed. Input from various remote sources leads to stronger variations in CO mixing ratio. However, all sectors contribute with more than 10% to the observed distribution. Our interpretation of air mass transport history is confirmed by the results from CO$_2$. 

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Figure 11. Phase-space diagram illustrating the maximum absolute change in temperature ($\Delta T$) and potential temperature ($\Delta \Theta$) relative to the time of the measurement for April 2015. The colour code denotes the CO mixing ratio at the time of the measurement. (a) shows all background data, (b) only shows those data corresponding to the polar dome and (c) includes all data points outside the polar dome. To separate the different regions the tracer derived polar dome boundaries are used (see Tab. 2).

Based on results from the phase space diagrams, we further analyzed the trajectories of the individual clusters. This allows for a more detailed analysis of the physical processes along the trajectory. We compare the two most dominant sectors for the April 2015 measurements in Figs. 12a-d and 13a-d for air masses inside and outside the polar dome, respectively. Sector 1 is mostly dominated by air masses confined to the central Arctic at all altitude levels (see Figs. 12a and b). The air masses show a weak descent during the 10 day period before the measurements but experience a very pronounced decrease in temperature and potential temperature (see Figs. 12c and d, median trajectory cluster temperature). In contrast, air masses outside the polar
domed, dominated by sector 4, contain contributions from different air streams. Air masses originate at different altitudes in the central Arctic, at low levels over the Pacific Ocean and from the upper troposphere over Asia. Air masses in this cluster are characterized by a significant increase in median temperature and decrease in median potential temperature indicating a descending trend, which is confirmed by decreasing median pressure over the time of travel. Low-level transport over the Pacific
Figure 13. (a) Trajectories of the most dominant sector 4 for air masses outside the polar dome. The color code represents the pressure along the trajectories. (b) The same trajectories as in (a) as a function of pressure and latitude, color coded by potential temperature. In both figures (a) and (b) black circles denote the initialisation point of the trajectory along the flight track. The black open squares show the position of the trajectory 10 days back in time. Figures (c) and (d) show a vertical cross section of the trajectory evolution over the 10 days of travel with the color code denoting the temperature (c) and potential temperature (d). The black line marks the median pressure of the trajectory cluster at the individual time steps and the grey line indicates the median temperature and median potential temperature, respectively. Note that in all figures only every 20th trajectory is plotted for figure clarity.

is associated with a low-pressure system over Alaska. Those air masses arrive at the polar dome boundary in the measurement region after experiencing a week net cooling over Alaska.

We conclude that air masses within the aged summertime polar dome are mostly confined to the boundary layer while they experienced a week diabatic warming due to insolation in July 2014 during NETCARE. In the mixing region and outside the polar dome diabatic cooling and a continuous descent is observed. Within the polar dome in April 2015 during NETCARE mostly near-surface processes (diabatic cooling due to the flow over cold surfaces) dominate the recent transport history of air masses in the lower polar dome. Air masses in the upper polar dome experience a very slow descent induced by radiative
Figure 14. Probability density functions (PDFs) of all CO (a and c) and CO₂ (b and d) background measurements during July 2014 (upper panel) and April 2015 (lower panel). The median latitude and maximum potential temperature coordinates along the 10 day back trajectories were used for the separation between inside and outside the polar dome using the tracer derived polar dome boundaries (see Tab. 2).

Cooling. Outside the polar dome, air masses mostly arrive at higher potential temperatures in the Arctic and experience a continuous slow descent with increasing temperatures but only week diabatic cooling.
5.4 **Chemical properties of the transport regimes**

5.4.1 **Trace gases**

Using the tracer-derived polar dome boundary we compare the composition of air masses within the polar dome region and the surrounding (see region (Tab. 2 for further details on the boundary values). The comparison is). We make this comparison based on probability density functions (PDFs) of measured trace gases CO and CO$_2$ for July 2014 (see Figs. 4214a and b) and April 2015 (see Figs. 4214c and d).

For July 2014 the three different regions identified in Sec. 5.2 are confirmed by the respective PDFs for both species. The aged polar dome and the mixing region show a quite similar distribution for both species, except differences in the mode of the PDF, but and are well separated from data outside the polar dome area. Whereas the absolute CO value inside the aged polar dome and the mixing region is lower compared to the area outside the polar dome, the CO$_2$ average mixing ratio within the polar dome is higher compared to the surrounding, as summarized in Table 3. This finding can be explained by the seasonal cycle of these two species and their zonal gradients (see data from NOAA ground based measurements in Fig. 64b). The minimum of the seasonal cycle of CO$_2$ in the Arctic and the mid-latitudes is reached at the end of the summer, typically during September. However, the onset of carbon uptake by vegetation in the mid-latitudes starts earlier compared to the high Arctic where less vegetation is present. At the same time the overall burden of CO$_2$ in the Arctic lower troposphere is to a large extent controlled by transport processes (Fung et al., 1983; Parazoo et al., 2011; Barnes et al., 2016). Mid-latitude air with relatively lower CO$_2$ mixing ratios is transported to the high Arctic, in particular during the second phase of the campaign. But, as the polar dome acts as a transport barrier for those air masses, exchange of high Arctic lower tropospheric air with mid-latitude air is reduced leading to the observed PDF for CO$_2$. Under 24 h daylight conditions and with only a few inner Arctic sources of pollution, CO concentrations reach their minimum in the high Arctic in late summer. Air masses transported into the high Arctic from more southern regions are expected to have relatively higher CO mixing ratios due to the seasonal cycle of CO that has a stronger amplitude in the Arctic compared to mid-latitudes (see Fig. 64a). An increased mid-latitude influence in the second half of the campaign was dominated by more mid-latitude influence, those air masses enhance the tropospheric enhanced the CO burden compared to inside the polar dome area which is burden in the Arctic troposphere, whereas the inner Arctic air masses observed in the first half of the campaign were dominated by photochemically aged low CO air during July 2014.

We observed a strong link between the trace gas distributions and the observed change in the synoptic situation from a more high pressure controlled regime to a synoptically active regime and the trace gas distributions was observed during the two distinct campaign phases of NETCARE 2014. Based on the trajectory simulations, increased mid-latitude influence was observed which in turn influenced the general concentration level of the trace gases CO and CO$_2$ (see Sec. 22). The fraction of trajectory points outside the polar dome as a proxy for mid-latitude influence increased from 37% to 79% 10 days before the measurements. This in turn led to an increase in the level from supplement Fig. S6). CO increased from 77.9 ± 2.5 ppbv to 84.9 ± 4.7 ppbv. At the same time CO$_2$ decreased from 398.2 ± 1.0 ppmv to 393.8 ± 2.3 ppmv. Furthermore an enhanced variability of increased variability in CO and CO$_2$ is observed, indicating enhanced entrainment of polluted
mid-latitude air masses into the high Arctic. Part of the trajectories originating from outside the polar dome area pass the Northwestern Territories at low altitude, potentially within the boundary layer, where extensive biomass burning was observed occurred during the time of the measurements and before. Accordingly, increased aerosol concentrations during the second half of the campaign were reported by Burkart et al. (2017).

Using the polar dome boundary values listed in Tab. 2 to separate air masses within the polar dome from those outside Fig. 42, Fig. 14 shows probability density functions for all CO (c) and CO2 (d) background measurements during April 2015. Based on the PDFs, the difference in the tropospheric trace gas composition within and outside the polar dome region and the surrounding is clearly visible for CO but not as distinct as for the July 2014 data set for, but less distinct CO2. Average values for both species in the respective regions are also summarized in Tab. 3. In general, the distributions of CO and CO2 are much narrower within the polar dome region and CO mixing ratios tend to be higher within the polar dome. For CO2 the mean mixing ratio is quite similar within and outside the polar dome. However, a difference in the general distribution is observed. The reason for a less distinct separation between inside and outside the polar dome area is indicated by the seasonal cycles shown in Figs. 64a and b. CO2 concentration levels concentrations are on a plateau in spring, with reduced concentration changes with time accompanied by and reduced latitudinal gradients. The CO mixing ratio has already started to decrease at the time of the our measurements. However, the extent of the polar dome is much larger during the winter months and spring compared to the summer months. Hence more influence from northern mid-latitudes and thus in turn, and more mid-latitude pollution sources, are expected inside the dome areasince, CO rich air masses originate in cold regions of Eurasia and are able to reach the high Arctic lower troposphere where they are trapped during winter and early spring (Klonecki, 2003; Stohl, 2006; Jiao and Flanner, 2016). This leads to relatively larger large CO levels inside the polar dome compared to the region outside. Air In contrast, air masses from more southern mid-latitudes that are transported above the polar dome, have already experienced photochemical loss of CO in near their source region.
Figure 15. Normalized probability density functions for particles with diameters between 5 and 20 nm (a) and larger than 100 nm for July 2014 (b). The colour code represents the three different regions identified during the polar dome analysis. Panel (c) shows the distribution of the fraction of particles containing trimethylamine with respect to the total amount of particles measured by single particle mass spectrometry (Köllner et al., 2017) in the maximum potential temperature and median latitude coordinate system. Note that for the reason of particle spectra statistics the resolution of median latitude and maximum potential temperature interval is reduced compared to the refractory black carbon distribution (BC) shown in panel (d). The enhancement in BC within the polar dome is most probably due to fresh local pollution.

5.5 Transport regimes

5.4.1 Aerosol

In order to analyse the processes dominating the recent transport history of observed air masses during July 2014 and April 2015, we apply the phase space diagram introduced by Binder et al. (2017). This requires determining the maximum change in potential temperature ($\Delta \Theta$) as the difference between the potential temperature at the time of the measurement ($\Theta_0$) and the previous potential temperature minimum or maximum ($\Theta_{\text{min}}, \Theta_{\text{max}}$) along the trajectory. Depending on which difference is larger in absolute numbers, the air mass has either experienced diabatic heating ($\Delta \Theta = \Theta_0 - \Theta_{\text{min}}$) Having
defined the polar dome based on trace gas gradients allows for a more detailed study on aerosol associated with the different air masses. Efficient wet removal and less efficient transport from lower latitudes lead to generally low aerosol concentrations (Stohl, 2006; Engvall et al., 2008) in the Arctic lower troposphere during summer. This is consistent with results in Fig. 15a and b. Observations of elevated levels of accumulation mode particles ($N_{>100} > 0$) or cooling ($\Delta \Theta = (\Theta_0 - \Theta_{max}) < 0$). An analogue analysis is made for the absolute temperature to determine if an air parcel dominantly gained or lost temperature recently before the measurement. This analysis allows us to cluster the data into four categories shown in Figs. 13a d and 14a e. The changes of potential temperature and temperature along the trajectories, which is indicated by the clusters, can be associated with processes affecting the respective air mass. Sector 1 ($\Delta \Theta < 0$, $\Delta$ regions outside the polar dome and subsequent transport to the measurement region. In parallel, regions within the polar dome were characterized by $N_{>100}$ smaller than 100 T < 0) mainly contains air masses which experienced diabatic cooling, which indicates either thermal radiation, evaporation or low-level transport over snow or ice covered regions and thus cold surfaces. In sector 2 cm$^{-3}$. In contrast, number concentrations of ultrafine particles ($N_{5<20}$) showed occasionally larger values within the polar dome compared to outside ($\Delta \Theta > 0$ Fig. 15a), $\Delta T < 0$) air masses gained potential temperature which indicates an ascending air mass that is diabatically heated by for example solar radiation or condensation processes. Sector 3 ($\Delta \Theta > 0$, $\Delta T > 0$) includes those air masses that experienced both an increase in temperature and potential temperature probably due to solar insolation. Finally, sector 4 ($\Delta \Theta < 0$, $\Delta T > 0$) combines air masses that lost potential temperature and gained temperature during transport. Those air masses are diabatically cooled and thus experience a descent. We apply this clustering approach to identify differences between observations in the different regions (polar dome, outside polar dome, etc.) and whether we can connect trace gas mixing ratios to the dominant process in a specific region. The different regions were separated using the boundary values listed in Tab. 2, which are based on the tracer derived polar dome boundary..

Phase space diagram illustrating the maximum absolute change in temperature ($\Delta T$) and potential temperature ($\Delta \Theta$) relative to the time of the measurement for July 2014. The colour code denotes the mixing ratio at the time of the measurement. (a) shows all background data, (b) shows only those data corresponding to the aged polar dome, (c) shows the data points within the mixing region whereas (d) includes all data points outside the polar dome. To separate the different regions the tracer derived polar dome boundaries are used (see Tab. 2).

Phase space diagram illustrating the maximum absolute change in temperature ($\Delta T$) and potential temperature ($\Delta \Theta$) relative to the time of the measurement for April 2015. The colour code denotes the mixing ratio at the time of the measurement. (a) shows all background data, (b) only shows those data corresponding to the polar dome and (c) includes all data points outside the polar dome. To separate the different regions the tracer derived polar dome boundaries are used (see Tab. 2).

For July 2014, three regions are of particular interest, (1) the aged polar dome, (2) the mixing region and (3) the region outside the polar dome (see Figs. 13a d). Within the aged polar dome sector 3 dominates, thus solar insolation is of significant importance to heat the lowest levels. Air masses residing within the lowest altitude experience diabatic heating potentially resulting in a slow and shallow convective lift of the air masses prior to the time of the measurement. Equal contributions come from sectors 1 and 4, dominated by diabatic cooling either through descent, low level transport over cold surfaces or evaporation. Almost none of the air masses are in sector 2, thus a significant ascent of air masses within the aged polar dome.
hardly occurs. In contrast, outside the polar dome area sector 4 dominates and generally diabatic cooling occurs (sector 1 and 4). Thus, indicating the formation of ultrafine particles occurred within the polar dome region (Burkart et al., 2017). Exemplary for aerosol composition, particulate trimethylamine (measured by single particle mass spectrometry) can be associated with sources within the polar dome local surface near diabatic processes seem to mostly affect the air masses which also has an impact on the chemical composition. Observed mixing ratios are highest in the aged polar dome associated with aged Arctic air and negligible mid latitude influence. Outside, air masses have been transported into the Arctic and started to descent caused by radiative cooling. Associated mixing ratios of these air masses are significantly lower and can be attributed to more mid latitude regions. Within the dominating sector 4 air masses experience descent once they have reached the high Arctic at higher altitudes. This can be regarded as a typical transport pathway during the summer with a fast uplift of air masses at mid-latitudes within convective and frontal systems followed by a northward movement and finally a descent into the high Arctic lower troposphere. In between, air masses in the mixing region are even more dominated by sector 4. Within this mixing region trace gas concentrations still show dome-like characteristics thus low level processes seem to dominate over episodes of mid-latitude transport associated with air masses with relatively lower concentrations. Results for confirm the derived transport history for July 2014.

In April 2015 the picture is quite different (see Figs. 14a-c). Inside the polar dome diabatic cooling dominates, in particular sector 1. The reason is diabatic descent due to radiative cooling in the absence of sunlight. In addition low level transport over cold surfaces significantly contributes to transport into the high Arctic. Associated mixing ratios of show rather large values that can be explained by the accumulation of anthropogenic pollution from inner Arctic and high northern mid-latitude sources during the winter months. At that time chemistry is reduced in the absence of sunlight leading to an increase in the atmospheric lifetime. Air masses within the polar dome are quite efficiently isolated from any significant southern mid-latitude influence. In contrast, (Fig. 15c), consistent with results in Köllner et al. (2017) and Willis et al. (2017). In contrast, the abundance of refractory black carbon can be linked to pollution sources outside the polar dome and subsequent transport to the measurement region (Fig. 15d), (Schulz et al., 2019)). To conclude, the method introduced in this study is a useful tool to interpret Arctic aerosol observations in the context of transport processes and sources within and outside the polar dome the picture is more diverse. Sector 4 dominates, thus diabatically cooled air masses potentially associated with the tendency to descent are observed. Input from various remote sources leads to a stronger varying mixing ratio. However, all sectors contribute with more than 10% to the observed distribution. Also for the April 2015 measurements the derived transport history is confirmed by the results from sector 4.

6 Discussion

Jiao and Flanner (2016) used the maximum zonal mean latitudinal gradient of 500 hPa geopotential height in the Northern Hemisphere to assess the impact of changes in atmospheric transport and removal processes due to climate change on the aerosol distribution in the Arctic. They deduced the polar dome boundary between 40 and 50°N during January which is further south compared to our tracer derived values. However, their method does not account for the lower troposphere, which
is essential for the diabatic processes affecting transport into the Arctic. For January, Klonecki (2003) determined a mixing barrier in the lower troposphere at 60°N at longitudes between 60°W and 105°W for a short lived artificial tracer (7 days atmospheric lifetime) emitted in North America and Europe. During summer they reported that the strong mixing barrier moves north following the location of the Arctic front. This is in the range of our horizontal polar dome boundary for the respective spring (66.0 to 68.5°N) and summer (73.5°N) season. In comparison to the Arctic front location, our analysis seems to give a more northern boundary for both July 2014 and April 2015. Furthermore, Klonecki (2003) reported an increasing mid-latitude influence with increasing altitude, in particular above 4 km altitude, which supports our findings of the potential temperature boundary being below 300 K for both seasons, which corresponds to an altitude below 4 km. The increasing mid-latitude influence with altitude is also in line with the results from Stohl (2006), who also used the Arctic front as a marker for the polar dome boundary.

The most isolated regions of the polar dome, where air masses experience the longest residence times, span a bigger area during late spring (April 2015) compared to summer (July 2014). This is in good agreement with previous studies, since it is already known that the polar dome extent is much smaller during summer compared to the winter months and processes and transport pathways controlling the composition of the high Arctic lower troposphere differ between both seasons (Klonecki, 2003; Stohl, 2006; Law and Stohl, 2007; Engvall et al., 2008; Garrett et al., 2010; Fuelberg et al., 2010). Our measurements during spring confirm a larger extent of the dome compared to summer and rather represent winter conditions than summer.

Stohl (2006) further defined an Arctic age of air and concluded that this age increased with decreasing altitude from 3 days between 5-8 km to around 1 week near the surface during the winter season (maximum 10 days in the North America region). During the summer season the air in the lowest 100 m of the troposphere is even older with values of 13-17 days north of 75°N. For a rough estimate of the upper limit of a transport timescale for mid-latitude air travelling into the summer polar dome during NETCARE, we can estimate the time at which the average mixing ratio within the polar dome was last observed at mid-latitude ground based observatories (for example Mace Head, Ireland). For CO₂ this gives a transport time of around three weeks, which is in the order of magnitude of the summertime Arctic age of air in the lowermost troposphere reported by Stohl (2006). This estimate assumes a transport controlled mixing ratio in the Arctic lower troposphere, which is justified by studies from Fung et al. (1983), Parazoo et al. (2011) and Barnes et al. (2016). Stohl (2006) further report that the Arctic troposphere is flushed on the time scales of 1-2 weeks in winter whereas in summer the corresponding timescale is twice as long. Assuming a similar short Arctic age during the spring season this implies that above the polar dome air masses can be transported within days from mid-latitude regions to the Arctic troposphere. Tracer concentrations will be further homogenized along isentropic surfaces when diabatic processes are slow compared to transport timescales (Klonecki, 2003). This is evident in a layer of similar CO mixing ratios above the polar dome for the observed April 2015 measurements (see Fig. 7c). During the July 2014 measurements increased diabatic heating due to convective and boundary layer heating in mid-latitudes can lead to an uplift of air masses at in mid-latitudes and further transport into the Arctic, which prevents an isentropic distribution during summer as evident in Fig. 7a and b. Several studies analysed the transport of mid-latitude air masses into the Arctic troposphere above the polar dome along those pathways mentioned before, without specifying the extent of the polar dome in more detail (Fuelberg et al.,
Aerosol distribution as a function of maximum potential temperature and median latitude. The median and maximum values were derived from every 10-day trajectory calculated along the flight track. The grey-scale colour code represents the average particle concentration per bin. The coloured bin frame represent the three different regions identified in Sec. 5.2 (see Tab. 2): aged polar dome (blue), mixing region (green) and outside polar dome (red). (a) contains the distribution of aerosol with a diameter larger than 5 measured with a Condensation Particle Counter (CPC) and (b) contains measurements of aerosol between 20 and 100 obtained with a scanning mobility system (SMS) (see Burkart et al. (2017)).

Based on the 2014 data set the influence of the change in the weather regime on and levels was quantified. Mid-latitude air masses with enhanced and reduced concentrations most probably due to biomass burning emissions entered the high Arctic lower troposphere pushing the polar dome northward, which significantly changed the levels of the two trace gases in the measurement region (see Tab. 2). Burkart et al. (2017) also reported a significant change in the aerosol loading during the first and the second phase of the campaign.

Having defined the polar dome based on trace gas gradients now allows for a more detailed study of aerosol within the polar dome. Efficient wet removal and less efficient transport from lower latitudes lead to generally low aerosol concentrations (Stohol, 2006; Engvall et al., 2008), especially within the Arctic lower troposphere during summer. As evident in Fig. 15a and b, the lowest aerosol concentrations (Burkart et al., 2017) were observed within the aged polar dome which was derived based on the trace gas gradients. Interestingly Willis et al. (2017) found evidence of secondary aerosol formation events which can play an important role in growing nucleation mode particles into cloud condensation nuclei (CCN) active sizes in the clean summertime Arctic lower atmosphere within the polar dome. These events are indicated through enhanced concentrations within the polar dome region (blue squares) in the aerosol distributions shown here. Furthermore, the aerosol data support the previously discussed threefold structure of the high Arctic lower troposphere in the Resolute Bay area north of 75° N. This is indicated in Fig. 15 through different concentrations levels of the aerosol particles in the respective region. Köllner et al. (2017) also observed a threefold structure in the Arctic troposphere based on the analysis of single particle measurements. The known polar dome extent is further used by Willis et al. (2019) and Schulz et al. (2018) to study vertically varying source regions and chemical processing within the polar dome.

7 Summary and conclusion

In this study we defined the polar dome boundary based on tracer gradients. For In July 2014, our results indicate that the horizontal polar dome boundary was found to be at the located at a latitude of 73.5° N. In the vertical a threefold structure established with the strongest gradient being observed at the trace gas gradient observed in a potential temperature range between 299 and 303.5 K, separating air masses within the polar dome from those outside. A second weaker gradient was found at a potential temperature of 285 K. Below this potential temperature the region was denoted We identified the region below 285 K as the aged polar dome, with the highest degree of isolation and thus the longest residence time of air masses whereas

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above 285 K and 299 K a mixing region established. The mixing region still shows significant characteristics of the polar dome region and clearly separates from outside the polar dome (see Fig. 49a). In April 2015 the tracer-derived boundary between inside the polar dome and outside was determined to be polar dome boundary was between 66.0 and 68.5°N and in a potential temperature range of 283.5 to 287.5 K.

Using the tracer derived polar dome boundaries PDFs of and values inside and outside the polar dome clearly showed a difference in the distribution and also the absolute value of the distribution maximum. In the tracer tracer scatter plot the polar dome separates from the surrounding by a different slope (April 2015) or a narrow group of data points at end of the highest and lowest mixing ratios (July 2014). The PDFs and the scatter plots confirm the different air mass properties inside and outside of the polar dome.

The processes dominating the recent (10 day) transport history were analyzed using a phase-space diagram based on Binder et al. (2017). For air masses outside the polar dome, diabatic cooling and a temperature increase was prevalent in both seasons—spring and summer (see Fig. 10d and Fig 1c). The associated transport pathway starts at mid-latitudes where air masses are lifted in convective or frontal systems followed by further northward motion towards the high Arctic, before the descent starts and finally descent into the lower Arctic troposphere. Predominantly identified North America and East Asia are identified as the source region for this pathway by previous studies (Klonecki, 2003; Stohl, 2006). These source regions for our observations, Europe as the the major source region associated with this pathway (Klonecki, 2003; Stohl, 2006), are also indicated from Figs. 4 and 5, however our observations (Figs. S1 and S2); however, a more comprehensive study of the source regions is beyond the scope of this paper.

For spring and summer Furthermore, air masses within the polar dome separate based on their transport history. During spring air masses experience predominantly diabatic cooling and lose temperature which can be associated to low level transport over cold surfaces. During summer an efficient cooling mechanism is missing. In fact, already cold air masses within the polar dome potentially experience a weak heating thus leading to a conditionally unstable lower troposphere and potentially weak lifting. Diabatic cooling rates determined from the trajectories are in good agreement to with the range of 1 K per day (radiative cooling) to several degrees per day (contact with cold and mostly snow covered surface) reported for diabatic processes by Klonecki (2003).

Using our tracer derived polar dome boundaries, PDFs of CO and CO₂ concentrations inside and outside the polar dome clearly show a difference in the distribution and also in the absolute value of the distribution maximum. Tracer-tracer scatter plots indicate that the polar dome separates from the surrounding regions with a different slope (April 2015) or a narrow group of data points at the end of the highest CO₂ and lowest CO mixing ratios (July 2014). The PDFs and the scatter plots confirm different air mass properties inside and outside of the polar dome. For aerosol chemical composition, particles containing trimethylamine were enhanced inside the polar dome, indicating the influence of inner-Arctic sources whereas enhanced refractory black carbon concentrations outside the polar dome indicate remote sources of pollution.

We conclude that the different differing chemical composition within and outside the polar dome allows for a trace gas gradient based definition of the polar dome boundary. The phase space diagram Phase-space diagrams helped to cluster the air masses based on their differing heating and cooling rates. This gives further insight in the processes that control the re-
cent transport history of the air masses within and outside the polar dome. The polar dome boundary derived in this study has already been used to study the source regions and chemical composition of aerosol within the polar dome (Schulz et al., 2019; Willis et al., 2019). Additionally, mixing and exchange processes along the polar dome boundary triggered by synoptic disturbances can be studied to shed light on additional pathways of mid-latitude pollution that reaches the Arctic lower troposphere. A polar dome boundary derived from gradients of chemical tracers can be further used for a quantification of the influence of inner Arctic and remote sources of pollution affecting the Arctic lower troposphere in a changing climate.


10 Author contributions. HB wrote the paper, with significant conceptual input from PH and DK and critical feedback from all co-authors. HB, FK, JS, MDW, JB, HS and WRL operated instruments in the field and analyzed resulting data. AAA analyzed flight data. WRL, JPDA and ABH designed the field experiment. DK ran LAGRANTO simulations, and HB analyzed the resulting data with input from DK.

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

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