Wintertime sub-arctic new particle formation from Kola Peninsula sulphur emissions

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Abstract. Metallurgical industry in Kola Peninsula, North-West Russia, form, after Norilsk. Siberia, the second largest source of air pollution in the Arctic and sub-Arctic domain. Sulphur dioxide (SO₂) emissions from the ore smelters are transported to wide areas, including Finnish Lapland. We performed investigations on concentrations of SO₂ and aerosol precursor vapours, aerosol and ion cluster size distributions together with chemical composition measurements of freshly formed clusters at the SMEAR I station in Finnish Lapland relatively close (~300 km) to the Kola Peninsula industrial sites during the winter 2019-2020. We show that highly concentrated SO₂ from smelter emissions is converted to sulphuric acid (H₂SO₄) with sufficient concentrations to drive new particle formation hundreds of kilometres downwind from the emission sources, even at very low solar radiation intensities. Observed new particle formation is primarily initiated by H₂SO₄-ammonia (negative-) ion-induced
nucleation. Particle growth to cloud condensation nuclei (CCN) sizes was concluded to result from sulphuric acid condensation.

However, air_mass advection had a large role in modifying aerosol size distributions, and other growth mechanisms and <u>condensation of other compounds</u> cannot be fully excluded. Our results demonstrate the dominance of SO₂ emissions in controlling winter-time aerosol and CCN concentrations in the subarctic region with <u>a</u> heavily polluting industry.

1 Introduction

- Sulphur dioxide (SO₂) is one of the main air pollutants, influencing the acidification of soils and freshwaters, defoliation and reduced vitality of forests, atmospheric aerosol formation, cloud properties, and adverse health effects by air pollutants. Anthropogenic SO₂ originates primarily from the combustion of fossil fuels at power plants, other manufacturing complexes and ships, as well as from the smelting of sulphur containing mineral ores. Because of the severe environmental and health effects by SO₂, efforts have been made in order to suppress its emissions to the atmosphere. While the global SO₂ emissions
- 30 <u>have not shown any rapid decay, emissions in e.g. OECD (Organisation for Economic Co-operation and Development)</u> countries have <u>decreased</u> significantly within last 3 decades (Solarin and Tiwari, 2020).

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The metallurgical industry with large_scale smelter complexes in the Kola Peninsula, North-West Russia, form the second largest source of air pollution in the Arctic and sub-Arctic region. Smelters emit large quantities of SO₂, metals and particulate matter to the atmosphere. These pollutants, especially SO₂, have large, impacts on both atmosphere and biosphere in the

- 60 surrounding area, including the eastern parts of Finnish and Norwegian Lapland. In the close proximity of industrial plants, these pollutants have literally destroyed ecosystems, creating "industrial deserts" (Paatero et al., 2008). Though emissions have significantly decreased from ca. 600 kilotons yr¹ in early 1990's (Tuovinen et al., 1993; Ekimov et al., 2001), partly because of the collapse of Soviet Union and related socio-economical changes in Russia, they are still high. Today, Kola peninsula SO₂ emissions are around 200 kilotons yr¹ (Barentz Observer, 2019), far higher than the SO₂ emissions of the whole Finland (37
- 65 kilotons yr¹ in 2017). Though vast, Kola emissions are still far behind the emissions of the World's number one SO₂ polluter, Norilsk (Krasnoyarsk Krai, Northern Siberia), with enormous 1.5 megatons yr¹ emission rates (Barentz Observer, 2019). Together with the few other smaller_scale industrial complexes, these smelters are almost the sole local sources of air pollution in the very sparsely_populated (sub-)Arctic Eurasia, and therefore understanding their role in atmospheric chemistry and physics is of great importance.

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In the atmosphere, SO₂ can be photochemically oxidized to sulphuric acid (H₂SO₄) in the gas phase. While most of the atmospheric sulfate is formed from SO₂ in a liquid phase in cloud droplets, which may evaporate leading to sulfate aerosol production or precipitate as acid rain with very high concentrations of SO₂ in the Kola Beninsula area also high production rates of gas-phase H₂SO₄ are expected. The H₂SO₄ vapour can, in turn, contribute to atmospheric new particle formation (NPF)
via nucleation and subsequent particle growth even up to sizes of cloud condensation nuclei (CCN) by further condensation of H₂SO₄ and potentially some other vapours (e.g. Weber et al., 1995; Kirkby et al., 2011; Jokinen et al., 2018). Atmospheric CCN formation globally (Merikanto et al., 2009; Gordon et al., 2017). At high latitudes, the contribution of NPF has been estimated to be even larger, reaching >90% of the cloud level CCN in the high Arctic and approximately 70-80% in our study area, the

80 sub-Arctic zone of Northern Finland and North-Western Russia (Gordon et al., 2017).

Vehkamäki et al. (2004) were the first to report observations of NPF (> 8_nm diameter particles) at the Värriö SMEAR I field station in eastern Lapland, Finland, relatively close to the Kola Peninsula smelter complexes. Their results on the contribution of SO₂ pollution were not completely definitive, so that during the four years of measurements 15 out of the 147 observed NPF
events were concluded to be explained by SO₄ pollution plumes, Kyrö et al. (2014) recorded particle number size distributions down to 3 nm in diameter and showed that NPF is connected to high concentrations of SO₂. They observed NPF even during winter in almost dark conditions, indicating that during episodes of very high concentrations of SO₂, a sufficient fraction of it is converted to H₂SO₄ in the gas phase even in very low solar radiation levels to initiate NPF. However, to date, no reports on

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115 quantification of sulphuric acid concentrations by direct measurements, or detailed mechanisms and chemical compounds involved in NPF, in this area exist,

While observation of atmospheric NPF has been reported in hundreds of publications since the times of John Aitken (<u>Aitken</u>, 1900), the details, i.e. the dynamics and contributing compounds, of NPF have been experimentally resolved only in a limited

- number of atmospheric environments. Pioneering studies include the observations by Weber et al. (1995) on the connection, between sulphuric acid and atmospheric nucleation, and the first report on ion-induced nucleation and simultaneous detection of sulphuric acid anion clusters using a mass spectrometer by Eisele et al. (2006). Among the field work, several laboratory investigations by the same research groups probed the properties of sulphuric acid water and sulphuric acid ammonia water clusters and their potential role in new particle formation (Ball et al., 1999; Hanson and Eisele, 2002).
 - Later advances in understanding the *molecular* steps of nucleation and growth in the atmosphere include the discovery that iodic acid (HIO₃) is primarily responsible <u>for</u> nucleation and growth in coastal areas and in the vicinity of the Arctic sea ice (Sipilä et al., 2016; Baccarini et al., 2020). Jokinen et al. (2018) demonstrated that in coastal Antarctica, H₂SO₄ <u>originating</u> from <u>the</u> oxidation of dimethyl sulphide (DMS, emitted by pelagic phytoplankton) and ammonia (NH₃, from penguin colonies)
- 130 nucleate via <u>a</u> negative ion-induced mechanism, with sulfuric acid condensation accounting for most of the subsequent particle growth. Further observations on nucleation mechanisms indicate the key role of highly oxidized organic molecules HOM (Ehn et al., 2014) in NPF <u>during</u> the spring-summer time <u>in a</u> boreal forest <u>environment (e.g. Kulmala et al., 2013; Rose et al., 2018)</u> and in the mid-latitude continental free troposphere (Bianchi et al., 2016) in parallel with sulfuric acid ammonia nucleation (Bianchi et al., 2016; Schobesberger et al., 2015; Yan et al., 2018). Dimethyl amine has been found to contribute to <u>initial</u>
- 135 nucleation in polluted urban air (Yao et al., 2018). In addition to these, yet rare molecular_jevel atmospheric observations, several laboratory studies have investigated the details of these nucleation mechanisms (e.g. Kirkby et al., 2011; Almeida et al., 2013; Kürten et al., 2014; Kirkby et al., 2016). Recent laboratory studies that probed nucleation of nitric acid and ammonia suggest the mechanism may contribute to new particle formation and growth especially in the upper troposphere (Wang et al., 2020).
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 - Mass spectrometers (Junninen et al., 2010; Jokinen et al., 2012) and air ion spectrometers (Mirme and Mirme, 2013), have largely facilitated the recent progress in the field of atmospheric NPF. By utilizing them in conjunction with aerosol and meteorological observations, this work aims to shed light on the molecular steps of NPF resulting from (sub-)Arctic air pollution during wintertime. Investigations were carried out at the SMEAR I research station in Värriö strict nature reserve in

145 Finnish Lapland close to the industrial plants (most of them located approximately 300 km east from the station) of Kola Peninsula, north-west Russia, during the winter 2019 – 2020.

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2 Methods

2.1 Site and time of the study

- 160 Measurements were carried out at the Värriö SMEAR I research station (Hari et al., 1994) located in Värriö strict nature reserve, Finnish Lapland, in the vicinity (5 km) of Russian border (Fig. 1) (67°45′19″N 29°36′37″E). The station stands on a top of a hill (390 m a.s.l.), being surrounded by untouched pine and spruce forests, bogs, fells, small lakes and rivers. Several large smelter complexes are located ~300 km away from to the station on the Russian side of the border, while on the Finnish side no smelters or other large-scale energy intensive (polluting) industrial plants are located within a distance of several hundreds
- 165 of km_e the closest, relatively small coal burning plant is located 550 km away. The SMEAR I station was set up in 1991 for monitoring air pollution, especially sulphur dioxide (SO₂) originating from the Kola Peninsula smelters. In this work we present 4.5 months data from winter time, 1st November 2019 until 16th March 2020.

2.2 Instrumentation

Aerosol number size distribution between 3 and 750 nm was recorded by a twin differential mobility particle sizer (DMPS) [170 (Aalto et al., 1999), comprising Hauke-type differential mobility analyzers (lenghts 110 and 280 mm) and TSI-3776 and TSI-

3772 condensation particle counters (TSI Inc., Shoreview, MN, USA) as detectors. <u>The other DMPS malfunctioned during 9th</u> – 10th and 14th – 27th January, resulting in the loss of data on 3 – 10 nm particles.

Number size distribution of charged particles and molecular clusters between 0.8 and 40 nm was recorded by a Neutral Cluster and Air Ion Spectrometer (NAIS, Airel Ltd., Estonia, Mirme & Mirme, 2013).

Aerosol precursor vapour concentrations, H₂SO₄, methane sulphonic acid (MSA), HIO₃ and HOM were measured by a nitrate ion – Chemical Ionization - Atmospheric Pressure interface – Time-Of-Flight mass spectrometer (CI-APi-TOF, Jokinen et al., 2012), equipped from 26th January onwards with a switcher inlet, with which the instrument can switch between chemical

- 180 ionization (CI) operation mode and natural ion detection mode. <u>This instrument was calibrated in the CI-mode for sulphuric acid, as described by Kürten et al_v(2012). <u>A same calibration coefficient was used for reported MSA and HIO3</u>. However, the instrument was not fully operational at all the times during the measurement campaign and significant fraction of data (including all collected prior 25th December and a long period in January) was disregarded.</u>
- SO₂ was recorded with a TEI 43 i-TLE pulsed fluorescence analyzer, O₃ by a TEI 49 i photometric analyzer, and NO₈ by a TEI 42C TL chemiluminescence analyzer with photolytic NO₂-to-NO converter, all manufactured by Thermo Fischer Scientific (Franklin, MA, USA). The wind speed and direction as well as the air temperature were measured with a Vaisala WTX sensor 16 m above ground level.



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2.3 Nucleation rate calculation

Negative (-) and positive (+) ion-induced nucleation rates of 1.5 nm particles, $J^{\prime_{1,5}}$, were calculated assuming a steady-state between formation and loss of particles in the size range of 1.5 and 2.5 nm:

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$$J_{1.5}^{-/+} = \frac{dN_{1.5-2.5}^{-/+}}{dt} + \frac{GR_2}{\Delta d_p} N_{1.5-2.5}^{-/+} + CoagS N_{1.5-2.5}^{-/+} + k_{rec} N_{<1.5}^{+/-} N_{1.5-2.5}^{-/+}$$
(1)

where $N^{-t_{1,5,25}}$ is the total concentration of negative or positive ions in the size range between 1.5 and 2.5 nm, filtered using Matlab's Savitzky-Golay 2nd order filtering method to remove the instrument noise, k_{rec} is the recombination coefficient between negative and positive small ions which was here, approximated by a size_independent constant of 1.6·10⁻⁶ cm³ s⁻¹ (Tammet, 1995), $N^{+t_{c1,5}}$ is the concentration of positive or negative sub-1.5 nm cluster ions. GR_2 is the 2 nm particle growth rate, Δd_p is the width of the size interval for which the concentration is defined ($\Delta d_p = 2.5$ nm - 1.5 nm = 1 nm)₃ and CoagS is the coagulation sink of 2 nm particles to the pre-existing background aerosol population. CoagS was calculated from the following equation;

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$$CoagS = \sum_{i=1}^{n} K_{2nm,i} N_i$$

(2)

(3)

where N_i is the concentration of particles in the channel *i* of DMPS and <u> K_{2nni} is the</u> coagulation coefficients <u>between a 2-nm</u> particle and a particle in the size bin *j* calculated based on Seinfeld and Pandis (1998).

A<u>n a</u>ccurate determination of <u>the</u> particle growth rate for 2 nm particles from the size distribution is challenging, and therefore GR_2 was approximated by assuming irreversible sulphuric acid condensation <u>as the sole mechanism of growth similar</u> to <u>Jokinen et al. (2018)</u> and Beck et al. (2021<u>)</u>according to (Stolzenburg et al., 2020);

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$$GR_2 = 1.45 \cdot \left(2.68 \cdot \left(\frac{d_p}{\text{nm}}\right)^{-1.27} + 0.81\right) \cdot [\text{H}_2\text{SO}_4] \cdot 10^{-7} \text{molec.}^{-1} \text{ cm}^3$$

where the pre-factor 1.45 accounts for dipole - charge interaction in charged particle growth (Stolzenburg et al., 2020). The justification for this approach will be discussed later. More standard method for GR determination is to approximate *GR*₂ by the average growth rate of the formed particle population, including mainly particles grown far above the 2 nm size, during a few hours starting from the beginning of the event as demonstrated in Figure 2A. This method leads to average GR of 4.5 nm h⁻¹. However, this approach neglects the effect of air_mass advection which, as will be discussed later, may largely determine the time development of the size distribution and thus also the apparent growth.

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Rather than the average GR of the whole particle population, the 50-% appearance time method (Lehtipalo et al. 2014) could be used to estimate the growth rate of nucleating clusters in the size range of 1.3 – 2.7 nm (Figure 2B). Here, the cluster appearance time in each size channel represent the time when cluster concentration reaches 50% of its maximum concentration during the event. Growth rate can be assessed from the cluster diameter vs. appearance time curve (black line) resulting in ~0.35 nm h⁻¹ between 10:10-11:30 and in ca. 1.8 nm h⁻¹ growth during 11:30-11:50 and the average growth rate of ~0.9 nm h⁻¹ during the period from 10:10-11:50. Drawback of this analysis is that the GR cannot be obtained for the period where concentration has passed the 50%-threshold or the period of decaying concentration. Furthermore, temporal GR cannot be properly obtained due to fluctuations in the data. Nevertheless, above mentioned values of 0.35 – 1.8 nm h⁻¹ can be compared to those obtained from Eq. 3, which yields maximum GR during the example day (29th January 2020) of 0.51 nm h⁻¹ around noon. Comparison to 1.8 nm h⁻¹ obtained from the 50%-appearance time method slightly before noon leads to a factor of ~3.5, difference in *GR*₂ on that day, which, in turn, is reflected in 22% difference in the calculated nucleation rate (Eq. 1). Effect of different determination of GR are visualized in Figure S1. To conclude, ion-induced nucleation rate calculation is not very sensitive to *GR*₂ because ion-ion recombination term (Eq. 1) dominates the loss in our conditions.

2.4 Sulphuric acid proxy calculation

Because of significant gaps in the measured data, [H₂SO₄] was also calculated using a proxy developed by Dada et al. (2020). <u>This proxy takes into account the oxidation of SO₂ to H₂SO₄ both by OH (estimated from the global radiation intensity) and stabilized Criegee Intermediates (estimated from monoterpene and ozone concentrations; Sipilä et al., 2014), as well as losses</u>

- 275 of, H₂SO₄ by dimerization (negligible in observed concentrations) and condensation onto pre-existing aerosol particles (the primary loss term). Unfortunately, there are no VOC measurements available at SMEAR I, but because the data were collected during winter well outside of the growth period, we assumed the monoterpene concentration to be zero. Global radiation measurements showed unexplained fluctuations (maybe caused by low solar zenith angles or freezing of the sensor) during the measurement period, and therefore we used UVB radiation and the relation between UVB and global radiation determined by
- 280 Dada et al. (2020). <u>During the times when CI-APi-TOF was operational, the agreement between the measured and calculated concentrations was good (Figure S2) with mean concentrations agreeing within 8%. Obtained correlation coefficient was R = 0.790 and coefficient of determination R² = 0.624.</u>

2.5 Trajectory analyses

285 Trajectories were calculated by using the Hybrid Single-Particle Lagrangian Integrated Trajectory model HYSPLIT (Stein et al., 2015) with GFS 0.25 degree meteorology as an input. We calculated 24-hour backward trajectories arriving at 50 and 250 meters above ground level, for the period 28.1.2020 00UTC to 30.1.2020 00 UTC, arriving every 6 hours. The trajectory calculations included mixing layer depth along the trajectory.

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3 Results and Discussion

315 3.1 New particle formation during the measurement period.

Figures <u>3a & 4a</u> depict the aerosol <u>number</u> size distribution between <u>3</u> and <u>700</u> nm, as recorded by the DMPS. Several new / particle formation events were <u>observed</u> during the measurement period. Clear <u>NPF</u> events with concentration of 4 -10 nm / particles, N_{4-10mm}. (Figures <u>3b & 4b</u>) exceeding 50 cm⁻³ are marked with grey shadings. Since the DMPS data on sub 10-nm / particles were missing from 9th -10th and 14th – 27th January N_{4-10mm} could not be derived for those time periods. **Still**, at least /

- 320 on 18th and 19th January we can see NPF that eventually produces particles larger than 10 nm in diameted. The observed NPF events, coincided mostly (ca, 50% of the cases) with clearly easterly, (~90°) winds (Figures 3c & 4c) and elevated H₂SO₄ concentrations (Figures 3d & 4d) calculated based on Dada et al. (2020). H₂SO₄ concentrations depend, besides condensation sink and UVB radiation, on the SO₂ concentration which is connected to both wind direction and air_mass origin. Clear examples of such, SO₂ pollution_adriven NPF events are, for example, those occurring during three consecutive, days on 10th 12th November, 2019, two days on 28th 29th January, 2020, and on 13th March, 2020. The data from 28th 29th January are
- discussed in detail <u>below</u>, wh<u>ereas the</u> data from the other two exemplary periods are presented in the Supplement (Figures S₃ S₁₁).
- Not all the NPF events occurred during the easterly winds. The events observed close to mid-winter, from early December until early January, occurred with westerly winds or during the transition of wind direction from west to east, in relatively low concentrations of SO₂ and in the virtual absence of day light and H₂SO₄. These low-H₂SO₄ mid-winter events were observed to start from sizes larger than <u>a few nm</u>, which means that nucleation did not take place in-situ in the surroundings of the SMEAR I <u>station</u>. Those particles <u>had</u> thus formed elsewhere and <u>were</u> transported to the measurement site by horizontal advection, or vertically down from above the mixed layer. <u>Compared with particles of a few nm in diameter, gas-phase H₂SO₄.</u>
- 335 is lost much more rapidly onto pre-existing particles after its production ceases, so the lack of H₂SO₄ is not excluding its primary role in NPF though not supporting such role either. Some of the mid-winter <u>NPF</u> events coincided with elevated SO₂ concentrations, suggesting that sulphuric acid may have be formed in the <u>measured</u> air mass earlier, while some, especially the relatively strong <u>NPF</u> event on December 3rd presented in the Supplement (Figures S9 S11), occurred in a virtual absence of SO₂ suggesting that sulphuric acid had, not been formed to a significant extent in that air mass. Currently, we thus cannot
- 340 explain the mechanism of NPF on that day. However, the NO₂ concentration was slightly (Figure SQ) elevated in the measured air mass, which might have been connected with the elevated source of particles. Nevertheless, most of the <u>NPF</u> events seemed to be connected with the presence of H₂SO₄.

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3.2 Case study 28th - 29th January 2020

To resolve the details of new particle formation and growth, we focus on 3 time periods with clearly occurring nucleation and particle growth. Here we show results from analysis of a 2-day period from 28th – 29th January 2020. To demonstrate that the discussed 2-day period is not only a unique observation, we represent data from time periods of 10th -12th November, 2019 (Figures S₃-S₃) and 13th March, 2020 (Figures S₅-S₃) in the Supplement. The data from the event on 3rd December, 2019, which differs from the overall picture, are also depicted in the Supplement (Figures S₂-S₁).

460 3.2.1 Meteorological situation and trace gas concentrations

Throughout the period of $28^{th} - 29^{th}$ January, the wind was blowing from the East (~50 - 150°) (Figure 5a). The ambient temperature ranged from -16°C to -28°C (Figure 5b). The sky was clear but, because of the low solar zenith angle (maximum 4.4° at noon 29th Jan.), the UVB radiation intensity needed for a photochemical H₂SO₄ formation reached only 35 mW m² (summertime UVB radiation intensity maxima at Värriö are >2000 mW m²). The HYSPLIT back trajectory calculations showed that air_masses arriving between January 28th at 6:00 and January 30th at 00:00 passed the industrial areas of the Montchegorsk region (Figure 6).

At_around 3:00 on 28th January coinciding with the change in the air_mass origin to the Montchegorsk-Kandalaksha region (Figure 6) (a few hours after the change of wind direction from west to east in the evening of 27th January), air pollutant levels
started to increase steeply (Figure 5d). During the course of the day, both SO₂ and NO₂ concentrations increased several orders of magnitude, with [SO₂] peaking at 27 ppb and [NO₂] at 7 ppb. Ozone (O₃) concentration declined from about 40 ppb to a 30 ppb range. To put the high level of SO₂ concentrations into some perspective, the highest concentration recorded at the Helsinki Metropolitan area was 8.4 ppb (24 µg m³, 1-hour average) in 2019 and the yearly-average concentration was about 0.2 ppb (~0.5 µg m³) (Helsinki Region Environmental Services Authority, 2020). The yearly-average SO₂ concentration, at the
SMEAR I station in 2019 was 1.1 ppb.

3.2.2 Aerosol precursors

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Despite the low UVB radiation, required for O₃ photolysis that initiates H₂SO₄ production via OH radical formation, H₂SO₄ concentration increased, from close to the lowest detection limit values of ~10⁵ molecules cm³ up to 8·10⁵ molecules cm³ during 28th and up to 1.5·10⁶ molecules cm³ on 29th (Figure 7c). Because the OH production rate must have been low, high
SO₂ concentration is a perquisite for the H₂SO₄ production during cold and dark winter months. Though stabilized Criegee Intermediates (sCI) formed in alkene ozonolysis can oxidize SO₂ to produce H₂SO₄ during summertime (Mauldin et al., 2012; Sipilä et al., 2014), alkene (terpene) emissions from the vegetation and thus the sCI production rate are negligible during the winter season. Proxy calculations agree well with the measured sulphuric acid concentration on 29th but show clearly higher

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values on 28th January. The cause of the disagreement on 28th January is probably the stable and shallow boundary layer. The temperature gradient close to the surface was almost +0.2 °C m¹₄ at noon on 28th (Figure 5b). Solar radiation from close the horizon is not penetrating efficiently inside the canopy, so the UVB measured above canopy and used in the proxy calculation does not reflect the situation at the ground level. Sulphuric acid produced above the canopy, on the other hand, does not mix downwards due to the strong temperature inversion and calm winds. On 29th January, the temperature gradient was absent or slightly negative, allowing the surface air to mix with the air above canopy.

Besides H₂SO₄, also minute signals of iodic acid (HIO₃) were, observed during the day (Figure <u>7</u>c). The exact production mechanism of HIO₃ remains globally unknown despite the emerging evidence on its critical role in new particle formation especially in the Arctic regions (Sipilä et al., 2016; Baccarini et al., 2020). Methane sulphonic acid (MSA), that has been observed in larger aerosol particles (Beck et al., 2021) and that could potentially also contribute to NPF, hardly exceeds the

detection threshold. This was expected since MSA originates from dimethyl sulphide (DMS) photo-oxidation. DMS ends up into the air mainly from the metabolism of pelagic phytoplankton during summer months, not during the dark winter. No other condensable vapours, such as HOM which dominate the new particle growth in the summer-time boreal forest environment

(Ehn et al., 2014), were observed during this case study period or during other periods depicted in Supplement.

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3.2.3 New particle formation

530 Ion size distribution

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Figures 7a and 7b show the NPF events on 28th and 29th January, as observed by the NAIS operated in the ion mode. Omnipresent small, < 1.5 nm ions are continuously produced by the galactic cosmic radiation, terrestrial gamma radiation and gas phase radon decay. Approximately at 11:00 on 28th January, coinciding with the increase of the H₂SO₄ concentration, small negative cluster ions started to grow, which is seen as small increases in the ~1.5th 2 nm negative ion concentration. During their growth beyond ~2 nm in diameter, those clusters were neutralized by collisions with positively charged ions so that they disappeared from the spectrum. They still obviously continued their growth, as charged particles reappeared in the spectrum after reaching some 5 nm in diameter when diffusion charging becomes effective enough; an equilibrium charging state for 2 nm particles is 0.8% while 5 nm particles are charged with an efficiency of 2.3% and out of 20 nm particles 11% are negatively charged (Wiedensohler et al., 2012). Opposite to negative ions, positive cluster ions did not grow in size. Larger, > 5nm positive particles (charged by diffusion charging during the course of their growth) grew similarly to negative particles. On 29th January, with clearly higher H₂SO₄ concentrations, the appearance of >1.5 nm negative clusters was more pronounced.

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suggesting higher rates of nucleation and critical role H_2SO_4 in the initial steps of NPF. Positive cluster ions were again only

bystanders and did not contribute to nucleation. This observation suggests negative ion-induced nucleation as the primary

pathway to new particle formation similar to H₂SO₄ - NH₃ (- H₂O) ion-induced nucleation observed by Jokinen et al. (2018)

in Antarctica and Kirkby et al. (2011) in CERN CLOUD chamber experiments. However, due to lack of information on neutral $\sim 1.5 - 3$ nm cluster concentrations, this observation alone does not exclude parallel neutral nucleation mechanisms.

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Nucleation rates

Even though weak growth of the small negative ions around noon on 28th January is visually observable in Figure 7a, the concentration of clusters in the 1.5_x2.5 nm size range (N^{-1,5,2,5}) is hardly distinguishable from the noise (Figure 7c). Nucleation rate, calculated using filtered concentration data, only slightly exceeded the baseline (caused by presence of minute, almost omnipresent signal from ion clusters extending above 1.5 nm but which is not connected to sulphuric acid nucleation), being approximately 0.005 cm⁻³ s⁻¹ with <u>a</u> high relative uncertainty (Figure 7d). On 29th, with <u>a</u> 2.3-fold sulphuric acid concentration, the concentration of 1.5_x2.5 nm negative clusters was well above the instrument noise reaching 20 cm⁻³ s⁻¹ around the noon. The nucleation rate peaked at 0.067 cm⁻³ s⁻¹. The ambient temperatures during nucleation (~ noon) were close identical, around

585 -22°C, in both days and therefore they can be directly compared. An approximately 10-fold difference in the nucleation rate between the two days accompanied by a factor of 2.3 difference in the sulphuric acid concentration is in line with the results from the CLOUD-chamber experiment on sulphuric acid – ammonia – water nucleation (Kirkby et al., 2011). The so-called "slope" that approximately (not exactly in real atmospheric situations) equals to the number of sulphuric acid molecules in the critical cluster (Vehkamäki et al., 2012) is given as:

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 $Slope = \frac{d \log J_{1.5}}{d \log [H_2 SO_4]}$

and yields a value of 2.9 for the numbers discussed above. Though this value is subject to a significant uncertainty, it would agree with observations of Kirkby et al. (2011) and parameterizations by Dunne et al. (2016) which yield a "slope" of approximately 3 under, similar conditions as visualized in Figure 8. In the same figure, data from all days with clearly observable ion-induced nucleation are depicted. There, hourly average nucleation rates $J_{1.5}^{-}$, that exceed a threshold value of $J_{1.5}^{-} = 0.01 \text{ cm}^{-3} \text{ s}^{-1}$, are plotted against the concurrent calculated sulphuric acid concentration and air temperature. Calculated nucleation rates, J_{GCR} , represent the total nucleation rates (ion-induced plus neutral) in different temperature and annonia concentrations under influence of galactic cosmic radiation (GCR) producing ions with the fixed rated of 1.8 ion pairs cm⁻³ s⁻¹. Negative ion-induced nucleation, however, should be the dominant mechanism under these conditions (Kirkby et al., 2011) and results can be therefore compared. Our data is reasonably close to range predicted by parameterization considering that this simple calculation does not include airmass transportation, vertical mixing, terrestrial radiation sources, or any other realworld phenomena. Also, sources and concentration of ammonia in our study area are unknown.

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Cluster time series

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To confirm the role of sulphuric acid and to solve the contribution of ammonia in the nucleation process, we measured the negative ion cluster composition and signal intensity with the APi-TOF in <u>the</u> ion mode without chemical ionization. The time series of <u>the</u> most abundant clusters show the appearance of HSO₄⁻ - ion in the morning of 28th Jan together with increasing [H₂SO₄] accompanied with a decay of NO₃⁻ - ion which typically dominates the anion spectrum at low [H₂SO₄] and low [HIO₃]

540 globally (Figure 2f). Since H₂SO₄ is a stronger acid than HNO₃, proton transfers from H₂SO₄ to NO₃⁻ explain the observed behaviour when [H₂SO₄] started to rise. When [H₂SO₄] still increased during the course of the day, (NH₃)_m(H₂SO₄)_nHSO₄⁻ - clusters started to form. The cluster signals peaked at around the noon coinciding with the highest [H₂SO₄] and N⁻_{1.5.2.5} after which they started to decay. On 29th January, the same behaviour was observed, but with somewhat stronger cluster signals due to the higher [H₂SO₄].

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Cluster composition

To get more insight into the chemical composition of clusters, the ion-cluster mass spectrum was integrated over 4 hours (2 hours effective data collection due to switching between CI and ion-inlet). The resulting spectrum is presented in Figure 9 by means of a mass defect plot, where the mass-to-charge ratio (*m*/*z*, unit Th) corresponds – with only singly charged ion clusters present in the air – to the mass of the cluster (*m*, unit Da, equal to unified atomic mass unit, u). Mass defect is the mass difference (in Th or Da) between the exact mass of the cluster and the integer mass defined as the sum of nucleons in the atomic nuclei of the cluster; for example, the exact mass of a HSO₄ -ion that has 97 nucleons is 96.960103 Da and the mass defect is thus 0.039896 Da. The area of a dot is proportional to the logarithm of the observed signal intensity. In the mass defect plot each addition of a molecule or atom is represented by a vector. Addition of a positive mass defect NH₃ molecule leads to an increasing mass and a decreasing total mass defect, while an addition of a positive mass defect NH₃ molecule leads to an increasing total mass defect. Successive additions of certain molecules to an ion results in a straight line in the mass defect plot, so that different cluster formation pathways are readily distinguishable from the plot.

In Figure 9, the largest signals are associated with omnipresent nitrate ion and its cluster with nitric acid (NO₃⁻ and HNO₃·NO₃⁻). The test of the small (<180 Da) ions are mainly different sulphur species, with bisulphate ion partly clustered with nitric acid (HSO₄⁻ and HNO₃·HSO₄⁻) being most abundant. Other small sulphur ions present in the spectrum are SO₄⁻, SO₅⁻, HNO₃·SO₃⁻ and HNO₃·SO₄⁻. Deprotonated iodic acid (IO₃⁻) and its nitric acid cluster, (HNO₃·IO₃⁻) are also abundant. Despite multiple different types of these core ions, their initial growth is caused solely by the attachment of sulphuric acid molecules. We observed clusters with 1-4 H₂SO₄ molecules attached to the SO₄⁻ - ion, one H₂SO₄ molecule attached to the to SO₅⁻ and SO₃⁻ - ion, sume that the negative charge remains in the core

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ion. This is not necessarily true, but H_2SO_4 may lose a proton e.g. to IO_3 resulting in the composition of HIO_3 (H_2SO_4)_{n-1}· HSO_4 instead of $(H_2SO_4)_n$ · IO_3 . Furthermore, water, if present in the clusters, efficiently evaporates in the vacuum of the mass spectrometer and therefore information on the role of water in the cluster formation is lost.

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None of the above discussed clusters seem to adopt ammonia efficiently enough for their signals to exceed the detection threshold of the APi-TOF (mass dependent, ~10⁻³ to few 10⁻³ ions/second for 2 hour integration for m/z > 400 Th). Only clusters made solely of sulphuric acid with a bisulphate ion (HSO₄⁻) as a core seem to efficiently attach ammonia, resulting in the formation of $(NH_3)_{m'}$ (H₂SO₄)_n·HSO₄⁻ -clusters (n>=3). This sequential addition of NH₃ and H₂SO₄ has been shown to be an effective (ion-induced) cluster formation and growth mechanism in coastal Antarctica (Jokinen et al., 2018) as well as a

- 695 an effective (ion-induced) cluster formation and growth mechanism in coastal Antarctica (Jokinen et al., 2018) as well as a secondary pathway in the free troposphere (Bianchi et al., 2016) and in the spring/summer time southern Finland boreal forest (Yan et al., 2018).
- Our results on negative cluster composition can be compared to the results from the CLOUD experiment at -25°C for varying
 NH₃/H₂SO₄ ratio (Schobesberger et al., 2015). Based on that data, with the NH₃/H₂SO₄ ratio exceeding approximately 100, both cluster composition and nucleation rate saturate (Kirkby et al., 2011), and become unaffected by further increases of the NH₃ concentration. In those conditions, a cluster comprising 3 molecules of sulphuric acid on a bisulphate ion, (NH₃)_n. (H₂SO₄)₃·HSO₄⁻, contains on average approximately n~1 molecules of ammonia, whereas a cluster composed of 4 molecules of sulphuric acid and a bisulphate ion, (NH₃)_n. (H₂SO₄)₄·HSO₄⁻, carries on average approximately n~1.5 NH₃ molecules
 (Schobesberger et al., 2015). In our case (Figure Q), corresponding average ammonia numbers were n~0.4 and n~0.8 for (NH₃)_n: (H₂SO₄)₃·HSO₄⁻ and (NH₃)_n(H₂SO₄)₃·HSO₄⁻ and (NH₃)_n(H₂SO₄)₄·HSO₄⁻, respectively, which would suggest that the NH₃/H₂SO₄ ratio in our case was well below 100, and likely below 10 (Schobesberger et al., 2015). If true, that would indicate an ammonia concentration
- of the order of ~10⁷ molecules cm³, or ~1 pptv. <u>However, cluster fragmentation inside the mass spectrometer can be totally</u> different between our experiment and Schobesberger et al. (2015) study and therefore any conclusions on ammonia concentration should not be drawn. <u>Nevertheless, if NH₃/H₂SO₄ – ratio would be low, that would mean that the system is not</u> saturated with respect to NH₃ and <u>that</u> the nucleation rate should therefore be sensitive to both H₂SO₄ *and* NH₃ similar to
- Jokinen et al. (2018). This together with unknown effects of cluster fragmentation, highlight the importance of understanding NH₃ sources, transportation and atmospheric mixing ratios down to ppt levels for a proper description of new particle formation also in the subarctic region. Unfortunately, NH₃ concentrations in the range of 1 pptv are not (reliably) detectable with any present-day measurement technology.

The present analysis shows that sulphuric acid – ammonia ion-induced nucleation can trigger new particle formation in the winter time sub-arctic / boreal environment with a high level of anthropogenic SO₂ pollution but a low UV-radiation intensity. Data on neutral 1.5_{2} nm particles are not available, so that neutral nucleation rates could not be derived. However, based on

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- 740 all the evidence obtained from field (mainly Jokinen et al., 2018) and especially from CLOUD experiments (Kirkby et al., 2011; Schobesberger et al., 2015), in the absence of significant amounts of compounds other than H₂SO₄ and NH₂ and with the nucleation rates being below the ion pair production rate (typically 2-4 ion pairs cm⁻³ s⁻¹ in the Earth's surface layer), ion induced nucleation seems to dominate over the neutral one. In our case, HOM were below the detection limit, and amines, if important, would appear also in the anion spectrum in H₂SO₄ clusters. HIO₃ and MSA were present, but significant neutral homogeneous nucleation of HIO₃ would require ~100-fold concentration of it compared to what was measured here (Sipilä et Compared to what was measured here (Sipilä
- al., 2016).

The observation of clusters containing IO_3^- or HIO₃ together with H_2SO_4 is, however, highly interesting. HIO₃ has been shown to nucleate homogeneously, and also mixed clusters containing both HIO₃ and H_2SO_4 have been reported from the Alps (Frege et al., 2017), the Atlantic coast (Sipilä et al., 2016) and the Arctic (Beck et al., 2021). If the SO₂ rich pollution plumes from

750 et al., 2017), the Atlantic coast (Sipilä et al., 2016) and the Arctic (Beck et al., 2021). If the SO₂ rich pollution plumes from the smelters are advected to iodine source areas (arctic ocean and especially sea ice zone as well as macroalgae rich coasts) or vice versa, this mixed mechanism may become important.

3.2.4 Particle growth and relevance for CCN-concentrations

Based on the above analysis, particle nucleation is clearly driven by sulphuric acid and ammonia, with the nucleation rate
being most probably sensitive to concentrations of both of these vapours. But how do the freshly nucleated clusters grow? Assuming irreversible condensation, even the peak sulphuric acid concentration of 1.5·10⁶ molecules cm⁻³ can explain only a small fraction of the observed growth rate. Consistent with an earlier report on wintertime particle growth rates at Värriö (Kyrö et al., 2014), the apparent average growth rate on 29th January was approximately 4.5 nm h⁻¹ (Figure 2), which, based on Stoltzenburg et al. (2020)₂ would require a steady concentration of ca. 2.6·10⁷ molecules cm⁻³ throughout the growth process which continues long after the sunset and the decay of [H₂SO₄]. Obviously, there are two possible explanations for this disagreement; either sulphuric acid is not responsible for most of the growth, or the air is not homogenous and the apparent growth is caused by the air mass advection.

Besides sulphuric acid, <u>the</u>only condensable vapours detected are MSA and HIO₃ (and NH₃). However, their concentrations are clearly lower than that of sulphuric acid, and therefore condensation of those in a homogeneous air_mass cannot explain the apparent growth either. It could be speculated that compounds not recorded by CI-APi-TOF, such as SO₂ or some <u>less</u> <u>oxidized</u> volatile or semi-volatile organic compounds, (S)VOC, <u>condense or</u> react in particle phase forming low volatile compounds thereby contributing to growth <u>(Stolzenburg et al., 2018)</u>, but a <u>complete absence of highly oxidized compounds</u> <u>does not support, though not fully exclude either, the presence of less oxidized compounds at a high abundancy. The NO₂</u>

770 concentration was moderate, up to 7 ppb, and therefore also nitric acid concentrations were likely insufficient to have a measurable effect on <u>the</u> growth (Wang et al., 2020). However, <u>the</u> temperature was low during the studied time period and

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therefore HNO3 or some other semi-volatile compound could have contributed to the growth, provided that such compounds, were present. Ammonia was detected in small ion clusters but its contribution to the particle volume concentration, assuming that the measured cluster NH₃/H₂SO₄ ratio reflects the composition of Jarger particles, was marginal. Assuming the particle composition to be ammonium bisulphate, i.e. NH₃/H₂SO₄ -ratio of unity, ammonia would contribute 17% to the particle volume and 5% to particle diameter growth rate.

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of particle populations.

The most plausible explanation for the observed growth is that the particle growth is driven by H_2SO_4 condensation but its concentration is not uniform over the source area. In that case, particles would nucleate and grow to their final sizes during the few hours of sunlight. Particles formed and grown in the close environment of emissions sources with high SO2 and thus H2SO4 805 concentrations grow to larger sizes than particles formed near the measurement site. Air mass advection would then transport particles through the dark hours, leading to a steadily increasing nucleation (and later Aitken) mode diameters at SMEAR I observed as apparent steady growth even through the night. Modelling efforts and measurement of chemical composition or hygroscopicity of growing modes would be required for an unambiguous explanation of the particle growth.

- 810 New particle formation in the sub-Arctic winter would be irrelevant if formed particles would not grow to sizes (above few tens of nm) where they can act as CCN. We did not measure CCN concentrations at different supersaturations, but the air masses arrived from the Murmansk-Kandalaksha region from about 03:00 onwards on 28th January, (Figure 6) contained elevated concentrations of Aitken and accumulation mode particles mainly in the size range of $\sim 20 - 500$ nm (Figure 10). New particle formation clearly increased the concentration of >3 nm particles, and also the concentration of particles larger than 50 815 nm showed an increase, especially on 29th January. The concentration of particles larger than 100nm was relatively constant and apparently unaffected by NPF during the times when these NPF events could be observed. Air mass advection and particle loss processes, however, naturally have an impact on measured concentrations and are largely responsible for the development
- 820 Figure 11 presents the average particle number size distribution during the about a one-week period of eastern winds (28th Jan - 3rd Feb 2020) when the two clear NPF events presented above occurred. Concentrations of particles in all the size classes were remarkably higher, even an order of magnitude for 10-200 nm particles, than the average concentrations during the preceding and succeeding time period with westerly winds. Concentrations during that one-week period were also clearly higher than the average concentrations between 1st November and 29th February, suggesting that new particle formation may 825 be a significant source of particles in the eastern air masses. However, primary emissions from the smelters and the surrounding cities would naturally show up in the size distribution plot as well. A more thorough analysis is needed to separate the roles of secondary NPF and primary emissions in the aerosol and CCN budgets. March, with almost constant NPF was excluded from this analysis since the lighting conditions in March differ significantly from those of early November - end of February.

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For an accurate assessment of contribution of secondary aerosol formation to CCN concentrations at SMEAR I or regionally, meteorological situation, including boundary layer dynamics, wet deposition of particles, etc. should be <u>considered</u>. However, our observations <u>on clearly elevated CCN-sized aerosol concentrations in eastern airmasses (Figs. 10 & 11)</u> point towards a <u>clear</u> contribution of Kola <u>Peninsula SO₂</u> emissions to winter<u>-time</u> CCN concentrations in the region.

4 Conclusions

 Winter-time new particle formation and growth was investigated at the SMEAR I station, in Värriö strict nature reserve, Finnish Eastern Lapland. Sulphur dioxide concentration in the airmasses arriving from Kola peninsula were often very high, occasionally over 30 ppb. In these high concentrations enough sulphuric acid formed even in very low solar radiation intensity to initiate new particle formation and growth.

New particle formation was observed mostly, but not solely, with easterly winds and in airmasses arriving from the direction of Kola peninsula smelters. Newly formed (4-10 nm, concentration > 50 cm⁻³) particles were observed in altogether 34 days
 between 1st November 2019 and <u>15th</u> March 2020 out of which ca. 60% were associated with eastern winds or with the period of wind direction change from ~west to east. Nucleation was observed in-situ at SMEAR I at H₂SO₄ concentration exceeding

ca. 1·10⁶ molec. cm³. These cases were identified based on the appearance of ~1.5-2 nm ion clusters. Other <u>NPF</u> events were observed as appearances of <u>a</u> few nm particles which gradually grew <u>in size</u> over time. Nucleation at SMEAR I was shown to proceed via <u>a</u> negative ion-induced sulphuric acid – ammonia (– water) channel which, based on prior understanding from

885 <u>laboratory experiments, can be hypothesized to dominate the NPF process at our site.</u> Closer to SO₂ emission sources where H₂SO₄ concentrations are likely remarkably higher, nucleation can proceed also via neutral channel and could, theoretically, involve compounds other than H₂SO₄, NH₃ and water.

Larger particles with a diameter of at least a few nm_observed at SMEAR I, were probably formed out of the immediate vicinity of the site and they had grown during the air_mass advection. Secondary aerosol formation from Kola emissions together with primary particle emissions impact the aerosol size distribution, clearly increasing the concentrations of particles in all the size classes, and therefore unavoidably also CCN concentrations. For better understanding of the contribution of Kola SO₂ emissions to local and regional CCN concentrations and upscaling our results to cover the whole (sub)-arctic Eurasia with vastly polluting industrial cities such as Norilsk, require more measurements. Those measurements should be complemented by CCN or cloud residual measurements – ideally in more than only one location (SMEAR I) around the Kola peninsula, Regional chemical transport and aerosol dynamic modeling would be necessary for thorough assessment. **Deleted:** Since the observed secondary particles are probably highly acidic, they are also highly hygroscopic, extending the CCN active fraction toward smaller particle sizes.

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Data availability. Mass spectrometer and air ion spectrometer data related to this article are available from Zenodo as well as upon request to the corresponding author. Rest of the data are available for download from https://smear.avaa.csc.fi/preview_

930 Supplement. The supplement related to this article is available online at:

Author contributions. MS designed the experiment, MS, NS, KN, TL, DK, SK, LB, TL, JL, PPA, PK, ES, PAR, and TJ prepared the instruments, performed calibrations, collected the data and processed the data, MS and NS analyzed the data, EMD calculated the back trajectories. MS wrote the manuscript. All authors contributed to the interpretation of data and

935 commented in manuscript.

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

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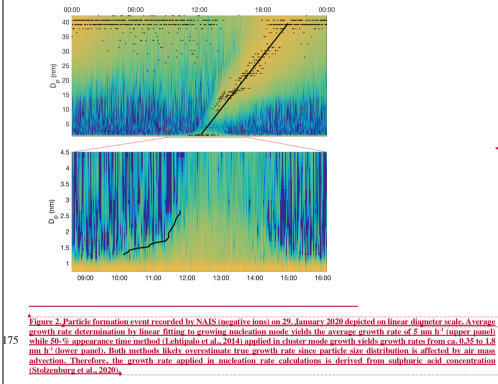
Figure 1. Map of the study area. Industrial cities of Nikel, Zapoljarnij, Monchegorsk, Kandalaksha have large<u>scale metal smelters</u> emitting vast quantities of SO₂ into the atmosphere. Kovdor and Olenegorsk mines produce nickel/iron ore, but have no smelter industry. Kirovsk and Apatity are phosphate mining and processing sites.

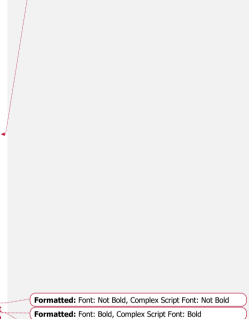
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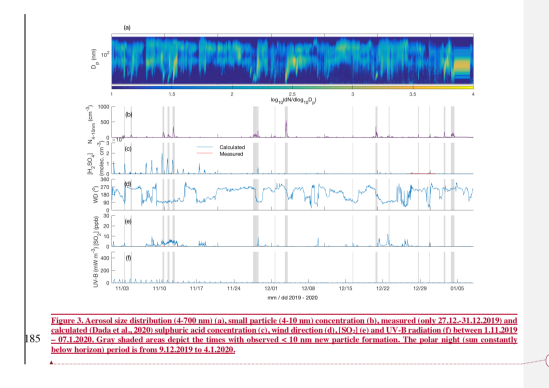


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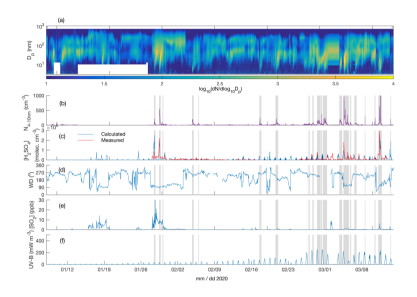
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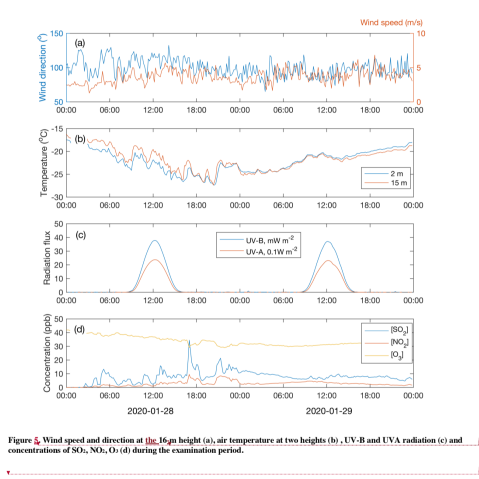


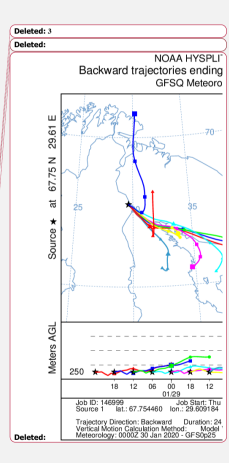
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190 Figure 4_Aerosol size distribution (4-700 nm) (a), small particle (4-10 nm) concentration (b), measured and calculated (Dada et al., 2020), sulphuric acid concentration (c), wind direction (d), [SO₂] (e) and UV-B radiation (f) between 8,1,2020 - 15,3.2020. Gray shaded areas depict the times with observed < 10 nm new particle formation.</p>

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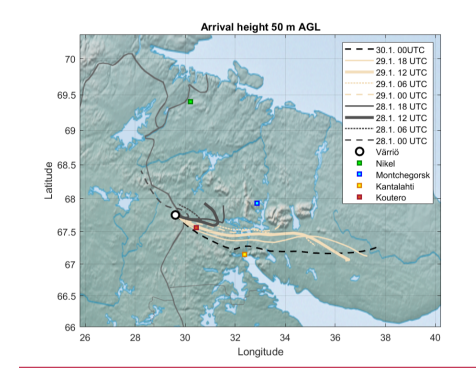




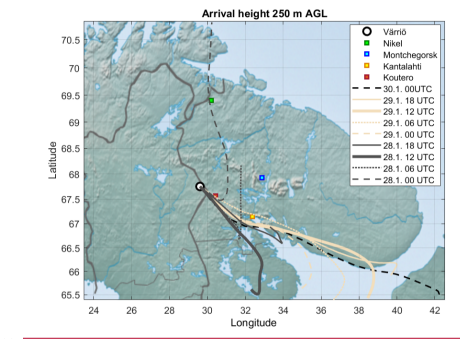


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Figure 6 24-hour back trajectories with an arrival height of 50 m (a) and 250 m (b) above ground level (AGL) and with an arrival, time between 28th January 00:00 and 30th January 00:00.

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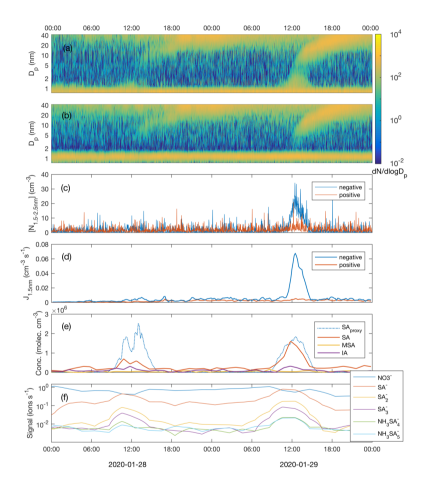
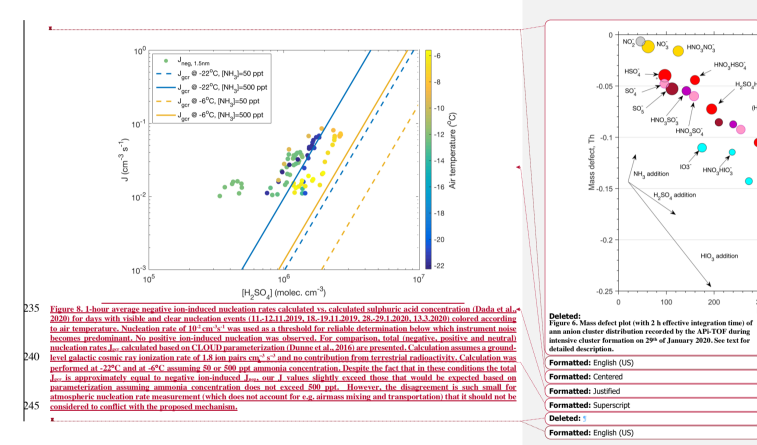
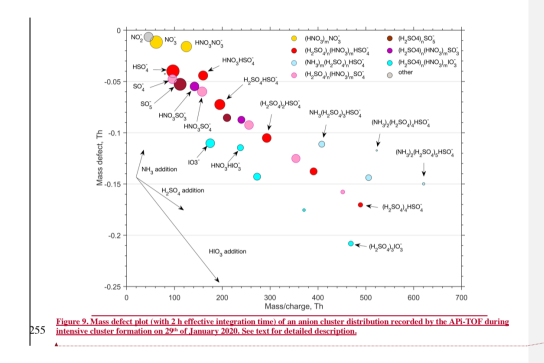


Figure 7, Size distribution of negative (a) and positive (b) clusters and particles, concentration of freshly nucleated, charged 1.5-2.5 nm clusters (c), formation rate of negative and positive 1.5 nm clusters (d), measured concentrations of sulphuric acid (HzSO₄), methane sulphonic acid (MSA) and iodic acid (HIO₃) as well as sulphuric acid concentration estimated by proxy calculation (e)₂ and the signal intensity of nucleating ion clusters with different composition (f) during the examination period.

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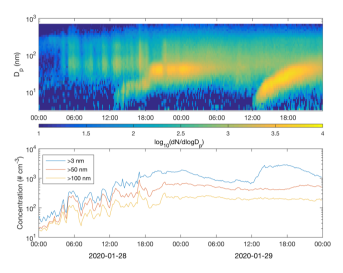


Figure 10, Particle number size distribution (a) and concentrations of particles larger than 3 nm, 50 nm and 100 nm (b) recorded by the DMPS. 260

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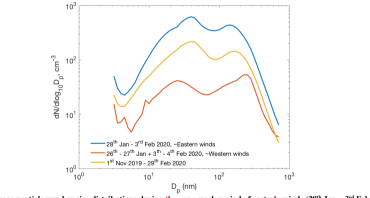


Figure 11. Average particle number size distributions during the week period of easterly winds (28th Jan – 3rd Feb 2020), during the preceding and succeeding time period with westerly winds, and average number size distribution between 1st November

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and 29th February. New particle formation in the eastern air_mass significantly increases the concentrations of particles in every 270 size class.

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