

Response to anonymous Referee #1 comments

The paper of Vodicka et al. aimed at elucidating particulate matter and their gaseous precursor sources by interpreting results of isotope analysis. The study is based on a year-round data set and, therefore, trying to discern seasonal differences and processes taking place in different seasons. The analytical approach to the measurement results is highly commendable where authors try to make sense of various relationships between the variables. By large the interpretations are justified although several of them are highly speculative and aiming to fit the data or simply shallow. Graphics of the paper is very well prepared and clear utilising three or four dimensions in 2D graph. The paper is certainly recommended for publication in Atmospheric Chemistry and Physics, but additional work is required by removing ambiguities, speculations and making the interpretations more coherent.

Response: Firstly, we would like to thank the referee for his positive criticism and valuable review which has enabled us to improve our paper. Based on the reviewer comments, we thought about different views of the issue and we also re-wrote some parts of the manuscript. Answers for reviewer's comments are following below.

Major comments

The major comment is the lack of coherent interpretation arising from unified approach to isotope effect and fractionation processes. Physico-chemical properties are the result of quantum mechanical effects with heavier isotopes (like C13 and N15) possessing lower vibrational energy levels and making stronger bonds and vice versa for lighter isotopes. Also, lighter isotope species have lower vapour pressures resulting in faster phase transitions. Condensation of isotope-light species would make the product of lower isotope ratio while evaporation would make it higher. Formation/evaporation of ammonium nitrate is, indeed, a reversible reaction, but not necessarily in equilibrium from isotope point of view because of competition effect with sulfate for ammonium. Particulate matter products, like ammonium sulfate or nitrate only consume a small fraction of their respective gaseous species pool (1-10%) with a possible exception of nitric acid (which is a product itself). Hence, isotope-heavy product does not necessarily involve heavy precursors, but instead isotope-heavy fraction of gaseous precursors due to the above considerations.

Response: Thank you for this comment. It provides some general information that were missing in our manuscript. Based on this, we decided to extend the Introduction chapter to the following paragraph: *"Isotopes are furthermore altered mainly by kinetic and/or equilibrium fractionation processes. Kinetic isotope effects (KIE) occur mainly during unidirectional (irreversible) reactions but also diffusion or during reversible reactions that are not yet at equilibrium (Gensch et al., 2014). Owing to KIE, reaction products (both gasses and particles) are depleted in the heavy isotope relatively to the reactants, and this effect is generally observed in organic compounds (Irei et al., 2006). If the partitioning between phases is caused by non-equilibrium processes (such as e.g. absorption), the isotopic fractionation is small and lower than that caused by chemical reactions (Rahn and Eiler, 2001). Equilibrium isotope effects occur in reversible chemical reactions or phase changes if the system is in equilibrium. Under such conditions, the heavier isotope is bound into the compounds where the total energy of the system is minimized and the most stable. Equilibrium effects are typical for inorganic species and usually temperature dependent."*

Particles of different size ranges do not induce isotope effects, but are rather a result of chemical transformations or being produced by different emission sources, e.g. traffic produces mainly nanoparticles, while solid fuel combustion produce wide range of particle depending on the size of the source (industrial scale versus domestic).

Response: We agree with the reviewer that just different particle size itself does not induce isotope effects on these particles, and we even do not say such statement in the paper so we are sorry if some part of text sounds so. We also agree with reviewer that in first round different emission sources have effect both on particle size and isotopic composition, which is valid especially for $\delta^{13}\text{C}$ values. But moreover, size of particles has an effect on different reactivity of these particles – e.g. compounds in small particles react more often than bigger one because their effective surface for reactions is larger. Submicron particles of accumulation mode also persist longer time in atmosphere so isotopic effects also take longer time than on coarse particles and may differ in the resulting isotope composition. The above implies that even if the same isotope effects occur on the particles, the indirect properties resulting from the particle size can lead to a different isotopic composition during particle chemical/physical transformations. And this is what we try to describe in the article.

For details, see answer for specific comment related to line 275 to effect of size on nitrogen isotope contents, and answer for specific comment related to line 366 due to carbon.

It would very useful to consider isotope mixing approach in interpreting the results which would eliminate unnecessary associations, e.g. $\delta^{13}\text{C}$ and EC (which is a relatively small fraction of TC). The authors, indeed, consider isotope mixing in few cases which is encouraged to do more frequently.

Response: Regarding to associations between $\delta^{13}\text{C}$ and EC we answered in specific comment related to line 454 (see below).

Regarding to multi-isotope approach, we extended Introduction chapter about examples of studies using multi isotope analyses – see following text:

“Recently, the multiple isotope approach was applied in several studies by using $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ measurements. Specifically, the $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ composition of aerosol (along with other supporting data) was used to identify the sources and processes on marine sites in Asia (Bikkina et al., 2016; Kunwar et al., 2016; Miyazaki et al., 2011; Xiao et al., 2018). Same isotopes were used to determine the contribution of biomass burning to organic aerosols in India (Boreddy et al., 2018) and in Tanzania (Mkoma et al., 2014), or to unravel the sources of aerosol contamination at Cuban rural and urban coastal sites (Morera-Gómez et al., 2018). These studies show the potential advantages of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ isotope ratios to characterize aerosol types and to reveal the underlying chemical processes that take place in them.”

Lastly, not all of the observations or measurement results should be strictly interpreted as some may be spurious or based on small subset of data and highly uncertain. Insignificant correlations should not even be noted with numbers, they are meaningless.

Response: Thank you for this comment. Actually, we had much bigger correlation matrix and we choose only correlations which were somehow interesting and also make sense for following interpretations. These value are summarized in Tables 2 and 3.

Regarding statistical significance/insignificance, this is determined based on p-value. Results are statistically significant if p-value is less than 0.05 in case of our study, and how it is used in case of many similar studies. However, the significance level of 0.05 is just a convention, and it is not entirely appropriate to omit the results of insignificant correlations just based on p-value. In fact, applying this discriminatory value is the subject of many disputations between statisticians, and using of p-value was even banned in some scientific journals (e.g. Siegfried, 2015, on-line).

In some cases, the difference between statistically significant and insignificant correlation can be very small - see for example the correlation of $\delta^{13}\text{C}$ with winter temperature (-0.35, $p=0.0328$ => significant) and autumn temperature (-0.33, $p=0.1063$ => insignificant) (Table 3). In other cases, this difference can be much greater, and comparison of such differences has meaning. We agree it is not necessary to comment insignificant correlations and in this sense we removed all insignificant correlations from text. However, we would like to keep these values in Tables 2 and 3 because it can be interesting for some readers, and it can be a good compromise of this situation.

Minor comments

Line 12. A study of stable carbon...

Response: Reworded

Line 15. 146 daily samples suggest 40% data coverage which is rather low for continuous sampling.

Response: We agree that the original formulation could be confusing. Sampling was performed every second day with 24-h time resolution and in this sense we changed this part of sentence from "*...collected on a daily basis at a rural background site...*" to "*...collected every two days with a 24 h sampling period at a rural background site...*".

Line 17. Autumn and spring were transition periods...(use past tense as the study refers to the past).

Response: Changed

Line 18. changing sources instead of different.

Response: Changed

Line 21. "Controls" can be used when processes are exactly known. "A comparative analysis with has revealed major associations which enlightened about certain processes affecting isotopic composition".

Response: The sentence was reformulated in sense of comment to the following text:

“A comparative analysis with water-soluble ions, organic carbon, elemental carbon, trace gases and meteorological parameters (mainly ambient temperature) has shown major associations with the isotopic compositions, which enlightened the affecting processes.”

Line 32. “...of nitrogen species.”, instead of “on nitrogen level”.

Response: Changed

Line 36. The winter event has occurred in prevailing southeast air masses.

Response: Reformulated

Line 43. “Aerosols have a strong impact on key processes in the atmosphere associated with climate change, air quality, rain patterns and visibility”.

Response: First sentence of introduction was reformulated in sense of comment. Thank you.

Line 47. Unique insights instead of information.

Response: Changed

Line 49. Is atmospheric history any different from physical and chemical processes in the atmosphere? What is meant specifically by history?

Response: We didn't mean different chemical and physical processes during history of atmosphere but different chemical origin of chemical compounds in atmosphere which is "signed" e.g. by changes of $\delta^{15}\text{N}$ of NO_3 in ice cores (see cited paper Dean et. al, 2014).

In this sense we changed end of sentence to: *“atmospheric composition in history”*.

Line 51. altered instead of influenced.

Response: Changed

Line 52. Delete "in case of nitrogen" and "in the case of carbon".

Response: These phrases were removed during rewriting of Introduction chapter.

Line 60. Americas or North&South America.

Response: Changed to “Americas”

Line 63. focusing on seasonality.

Response: Changed

Line 85. ...and to reveal undergoing chemical processes.

Response: Changed

Line 86. to broaden the approach over the European....

Response: Changed

Line 89. First study in the location or region? Surely authors must be certain about their location/station.

Response: We meant Central European region, so we changed word "*location*" to "*region*".

Line 95. ...observatory is a key station of the Czech..., focusing on air quality and environmental monitoring.

Response: Reformulated. Thank you.

Line 99. with very low traffic density.

Response: Changed

Line 110-112. Were 146 samples obtained as a result of continuous or strategic sampling? Unclear sampling strategy or low data coverage. Please explain. Perhaps "Some temporal gaps were caused by sampler maintenance or power outages resulting in 146 samples during a year-long study".

Response: We are sorry that our text was little bit confusing. Sampling was not made continuously day by day but every second day (it means 24 h of sampling followed by 24 h gap). Such kind of sampling was made based on strategic decision before starting of campaign. Even so we had three bigger gaps caused by sampler maintenance or power outages.

We reformulated sentences on lines 110-112 to make a text clearer. Instead of "*Aerosol samples (n = 146) were collected for 24 h every two days from September...*" we used "*Aerosol samples were collected two days for 24 h from September...*".

The sentence "*Some gaps in sampling were caused by outages and maintenance to the sampler.*" was changed to "*Some temporal gaps were caused by sampler maintenance or power outages resulting in 146 samples during the almost year-long study.*".

Line 113. sampled at a flow rate of 2.3 m³/h.

Response: Changed to: *"...operated at the flow rate at a 2.3 m³/h."*

Line 119. The PM1 mass was measured gravimetrically with a microbalance (...) in a controlled environment (...).

Response: The paragraph was rephrased and shortened in a sense of comment. Thank you.

Line 135. Thermo Electron Corp.

Response: Corrected

Line 136. ...for calculating TC, TN and their isotope values.

Response: Reformulated

Line 152. measured, not detected.

Response: Changed

Line 175. ...instrument response without filter exposure.

Response: Reformulated

Line 184. ...for the annual dataset (139) and separately for each season and winter event.

Response: Corrected

Line 186. ...for the correlation analysis as their distinctly high concentrations and isotopic values might have affected the results.

Response: Sentence was completed in a sense of comment. Thank you.

Line 193. Statistically, 146 sample may be sufficient, indeed, but specific season(s) may not be typical, unless known to be such from previous studies.

Response: Seasonal comparison with other years provide e.g. work of Mbengue et al. (2018) (also cited in our work), which is published 4 years survey of EC/OC together with other variables (e.g.

temperature) between years 2013-2016 directly at the Košetice station. It shows that a period from this work is not seasonally atypical during last years of observation.

Line 203. ..., but the relationship split during high concentration events due to divergent sources.

Response: Reformulated. Thank you!

Line 208. ...and characteristic of significant shift in chemical composition.

Response: End of sentence was changed in terms of comment.

Line 211. Use past tense as the study has been conducted in the past.

Response: Sentence changed to past tense

Line 209-211. Little difference does provide information on sources and quite contrary suggest that they were similar throughout the year in terms of BC production. The split contribution in each season may be different as suggested by isotope ratios, but overall the mixture of the sources seems to produce a steady trend.

Response: Station Košetice is a background site where the aerosol is more homogeneous than e.g. at an urban site, which can be reason for little seasonal differences of TC/TN ratio. However, previous studies from this station (Mbengue et al., 2018; Schwarz et al., 2016; Vodička et al., 2015) show that long-term concentrations of EC (and thus BC) are different in winter and summer, and sources are not similar throughout the year in terms of EC production. Different summer EC sources are consistent with a slightly higher summer TC/TN ratio than other seasons. In this sense, we changed the text to following:

“As seen in Table 1, the seasonal averages of TC/TN ratios fluctuate, but their medians have similar values for autumn, winter and spring. The summer TC/TN value is higher (3.45) and characteristic of a significant shift in chemical composition, which is in line with previous studies at the site (Schwarz et al., 2016). However, seasonal differences in the TC/TN ratios were not as large as those in other works (e.g., Agnihotri et al., 2011), and thus, this ratio itself did not provide much information about aerosol sources.”

Line 215. Do not report what was insignificant as it may mislead readers. “This result highlights significant shift in carbonaceous matter sources and corresponding isotope values in spring while during other seasons the sources were rather stable”.

Response: Thank you for this comment. Instead of original sentence we used above suggested one by the reviewer. We also deleted values of insignificant correlation coefficients in text.

Original sentence

“However, there is a statistically significant correlation for spring only (-0.54), while in other seasons, correlations are statistically insignificant (autumn: -0.29, winter: -0.11 and summer: 0.07). This result

shows that significant and related changes in the isotopic composition of nitrogen together with carbon occur especially in spring, while there are stable sources of particles during winter and summer.”
was changed to

“However, there is a statistically significant correlation for spring only (-0.54), while in other seasons, correlations are statistically insignificant. This result highlights a significant shift in the sources of carbonaceous aerosols and their isotope values in spring while the sources were rather stable during other seasons.”

Line 217. Lack of correlation during particular season is due to stability of sources while the variability between samples is similar in all seasons. Authors may look at the fractional variability of isotope ratios in each season as it seems that relative variability of $\delta^{13}\text{C}$ is a lot larger than $\delta^{15}\text{N}$.

Response: We agree with the review that lack of correlations during seasons is due to stability of sources in this period, and in this sense we also changed the sentence on line 215 (see previous comment).

In case of reviewer suggestion related to fractional variability of isotope ratios, we are not sure by its meaning because variability of $\delta^{15}\text{N}$ is larger than variability of $\delta^{13}\text{C}$.

Line 224. ...which increased by an average of 13permile.

Response: We changed original word "deviated" to "showed" instead of reviewer proposed "increase" because change is from 15 to 13‰. Whole sentence is following: *“The $\delta^{15}\text{N}$ values are stable in winter at approximately 15‰, with an exception of the winter Event, which showed by an average of 13‰.”*

Line 232. What is the purpose of the formula if not solved for fractions (which is impossible given one equation and at least two unknowns).

Response: The reason for showing this formula is to give readers an idea before discussion about contribution of different nitrogen compounds to $\delta^{15}\text{N}$ of TN and also to discussion about results presented in Fig. 3. However, because it is an equation with general information character, we decided to move this equation to the Introduction chapter.

Line 236. What does it mean similarly balanced if NO_3 was higher in winter?

Response: The original sentence was: *“...higher in winter, similarly balanced in spring and autumn, and very low in summer...”*. We wrote that similarly balanced were values in spring and autumn, not in spring and winter. However, we split this text to two following sentences to make this part of document clearer.

“...the highest in winter, and somewhat lower in spring and autumn. In summer when the dissociation of NH_4NO_3 plays an important role the NO_3^- content is very low and its nitrogen is partitioned from the aerosol phase to gas phase.”

Line 258. (a) changes in NO_x emissions

Response: Changed

Line 262. "Considering our study, it was most likely that all of the factors contributed to a certain extent to isotopic composition throughout the year".

Response: Original sentence was changed as follows: *"In our study, it is most likely that all these factors contributed, to a certain extent, to the nitrogen isotopic composition of NO₃⁻ throughout the year."*

Line 266. In summary,.... If enrichment of N15 occurs during lowest NO₃ contribution it can be inferred that NO₃ is depleted in N15. Is this inference consistent through the year?

Response: In our study, we observed the highest enrichment of $\delta^{15}\text{N}$ in TN during summer when NO₃ concentrations are lowest. However, during winter Event when NO₃ contribution was also on the lowest level, $\delta^{15}\text{N}$ in TN was contrariwise most depleted (see Figs.3 and 4). This Event shows that that exceptions may occur and we can't generalize. So, in our study, the inference, which you are proposing in comment, is not consistent throughout whole measurement campaign.

Line 275. Size fraction has no impact if the most of nitrogen containing particles reside in submicron range. Life-time has no impact either if coarse particles do not contain appreciable amount of nitrogen. If they do, what compounds that would be and how did they end up in coarse particles. If those compounds appeared in coarse particles by condensation then nitrogen was mainly concentrated on the surface and consequently coarse particles would be as reactive as fine ones.

Response: Nitrogen from NO₃ is contained in sufficient amounts both in fine and coarse fractions (e.g. Ondráček et al., 2011; Pakkanen, 1996; Schwarz et al., 2012), not only in submicron range. As summarized by e.g. Kundu et al. (2010), coarse mode contains predominantly non-volatile nitrogen in a form of NaNO₃ or Ca(NO₃)₂, whereas fine mode consists mainly from semi-volatile nitrogen from NH₄NO₃ and also in form of ammonium sulfate and bisulfate. If we have non-volatile nitrates in coarse fraction and predominantly NH₄NO₃ in fine fraction, where dissociation of NH₄NO₃ play an important role in enrichment of nitrogen, so it leads to some effect on the isotope composition depending on the particle size fraction. Yeatman et al. (2001) proposed presence of two different size-shift processes: dissociation/gas scavenging and dissolution/coagulation. Dissolution/coagulation processes appear to exhibit negative isotopic enrichment of nitrogen and shift both NH₄⁺ and NO₃ to the coarse mode, whereas dissociation/gas scavenging processes appear to exhibit positive enrichment factors. All this is supported also by the works of Mkoma et al. (2014) and Freyer (1991) who observed a higher enrichment of ¹⁵N in the fine fraction of the aerosol in comparison with coarse one. Last but not least, fine aerosol has larger particle surface/volume ratio than coarse one which can suggest higher reactivity of smaller particles.

Above arguments lead us to the fact that we have in this case the opposite view than reviewer.

Line 277. Aitken mode contributes negligibly to PM1 mass making this argument very weak. Unless authors can quantitatively prove it otherwise.

Response: Thank you for this notice. We made a mistake in this part. Instead of Aitken mode, we should write Accumulation mode there because this mode contributes by a main part to PM1 mass and also persist the longest time in the atmosphere. In text, we changed word "Aitken" to "accumulation".

Line 303. Not all of gaseous precursor mass is ending up in NH₄NO₃, but preferentially heavier part. Authors must consider kinetic fractionation, otherwise conclusions are biased or unfounded.

Response: It is clear that during incorporation of nitrogen from gas phase to aerosol phase play a role both equilibrium and kinetic fractionation. Equilibrium fractionation is related to bond stability of nitrogen isotope whereas kinetic fractionation is related to the "speed" of isotope. First time, nitrogen incorporation is probably driven by kinetic fractionation because lighter isotopes react faster, but later heavier isotopes form a more stable bonds during equilibrium fractionation. In fine fraction of aerosol, we have nitrates almost exclusively in a form of NH₄NO₃ which undergo to dissociation to NH₄⁺ and NO₃⁻ in water – this state is reversible and equilibrium fractionation is preferred in such system.

Nevertheless, Ciężka et al. (2016) suggested a possible kinetic exchange reactions between NH₃ and NH₄⁺ as one of three possible processes affecting nitrogen isotopic composition, especially for fossil fuels combustions during the heating season. Also Deng et al. (2018) reported the kinetic nitrogen fractionation factors between gaseous and aqueous ammonia with statement that, when the removal of degassed ammonia is not efficient, ammonia may dissolve back to the fluid, which may significantly shift the nitrogen isotope behavior from kinetic isotope fractionation toward equilibrium isotope fractionation. Indeed, all this suggests that kinetic fractionation is likely to affect the isotopic composition of fresh particles from combustion having lower δ¹⁵N than in spring or summer, and before they are affected by equilibrium fractionation.

Originally, on line 303, we did not present equilibrium fractionation as a dominant process, but we only compared known values of the isotopic composition of nitrogen oxides for different sources.

Since it is difficult to determine exact contribution of these fractionations to the final value of the heavier isotope in the aerosol, we do not discuss exactly, however, we added following two sentences to consider an influence of kinetic fractionation in a first steps of gas-aerosol transformation: *"Because of the only slight difference between above reported δ¹⁵N of nitrogen oxides and our δ¹⁵N of TN during maximal NO₃⁻ events, the isotope composition is probably influenced by the process of kinetic isotopic fractionation in fossil fuel combustion samples during heating season as referred by Ciężka et al. (2016) as one of three possible processes."*

Line 317. If ratio goes up during evaporation, NH₄NO₃ must have had lower ratio which makes sense for highly volatile compound. One could hypothesize that ammonium N15 ratio is the same in ammonium sulfate and ammonium nitrate, but as compounds are of different volatility that is unlikely, because volatile particulate compounds originate from lighter (more depleted) precursors than less volatile compounds which originate from heavier precursors.

Response: We agree with the reviewer that ammonium ¹⁵N ratio is NOT same in ammonium sulfate and ammonium nitrate and we didn't hypothesized opposite view in paper. On line 317 we wrote *'...the dissociation process of NH₄NO₃ can cause an increase in ¹⁵N in TN during a period of higher ambient temperatures...'* which is supposed to be okay, because evaporation of more volatile

ammonia from NH_4NO_3 comes more easily during the higher temperature and the lighter isotope is released into the gaseous phase with a higher probability.

Line 332. "Thus it can be considered..."

Response: Changed to "considered".

Line 342. more depleted, not "smaller".

Response: This part of text was removed during revisions.

Line 349. What does it mean "smaller"? Lower, more negative? That is not because of overlap, but source specific ratios which in case of organic carbon are largely negative.

Response: We are sorry for a confused formulation that may imply that we compare the negative values of $\delta^{13}\text{C}$ in comparison with $\delta^{15}\text{N}$. However, it is not so because we know that strongly negative $\delta^{13}\text{C}$ values originate from chosen carbon standard (PDB). Originally, we wanted to say that the range of TC $\delta^{13}\text{C}$ values is significantly smaller than a range of TN $\delta^{15}\text{N}$ values.

Based on this, we changed original sentence: "The $\delta^{13}\text{C}$ values are significantly smaller than those of $\delta^{15}\text{N}$..." to "The range of TC $\delta^{13}\text{C}$ values is significantly narrower than that of TN $\delta^{15}\text{N}$..."

Line 366. Wrong conclusion as mentioned in major comments. Fine particle sources are different from coarse particle sources. The ratio can only become more depleted in the atmosphere due to condensation of depleted precursors and even then condensation prefers heavier molecules, not lighter.

Response: The conclusion on line 366 that fine particles have lower $\delta^{13}\text{C}$ values than coarse particles was consistent with the Masalaite et al. (2015) and Skipitytè et al. (2016) studies referred in the same sentence on line 367, but is probably inappropriately formulated. We agree with reviewer that aerosol sizes itself cannot induce isotope effect and differences are caused e.g. by different aerosol sources. In this sense, we changed the sentence on line 366 from the original "...relatively low $\delta^{13}\text{C}$ values in our range (up to -28.9‰) are caused by not only sources of TC but also a fact that fine particles are more ^{13}C depleted in comparison with coarse particles (e.g., Masalaite et al., 2015; Skipitytè et al., 2016)." to the following: "relatively low $\delta^{13}\text{C}$ values in our range (up to -28.9‰) are caused because fine particles have lower $\delta^{13}\text{C}$ values in comparison with coarse particles probably due to different sources of TC. (e.g., Masalaite et al., 2015; Skipitytè et al., 2016)."

Line 389. "The aforementioned studies concluded that the isotope equilibrium exchange..."

Response: Changed.

Line 397. resulting relationship, not final.

Response: Changed

Line 401. Very narrow temperature range can produce unreliable relationships. The temperature range in this study is far more impressive.

Response: Thank you, we agree that temperature range is one of benefit of our study. (min. in winter -9.8°C to max. in summer $+25.5^{\circ}\text{C}$ results in $\Delta 35.3^{\circ}\text{C}$). It is visible that if we account data from whole year (and so we take the full temperature range) we have stronger correlation between $\delta^{15}\text{N}$ and temperature than just for individual seasons.

In cases of statistically significant seasonal correlations during autumn and spring we have following temperature ranges and correlations: autumn $\Delta T=15.8^{\circ}\text{C}$, $r=0.58$; spring $\Delta T=17.2^{\circ}\text{C}$, $r=0.52$. Pavuluri et al. (2010), whose work we compare, observed a strong correlation ($r^2 = 0.58$) for a temperature range of $\Delta T=6.1^{\circ}\text{C}$, which is a stronger correlation for a narrower temperature range than in our case. This gives an assumption that even during the narrower temperature range in the work of Pavuluri et al. (2010), we can get a relationship which is reliable for our comparison.

We did not make any revisions in MS related to this comment.

Line 413. What is the actual process of aging? Isn't it just the production of ammonium sulfate? Sure, production is two step: first bisulfate, than sulfate. It is obvious that decreasing molar ratio corresponds to lower nitrate, because ammonium nitrate can only be produced if at least bisulfate has been produced. When nitrate is not competing for ammonium due to higher temperature, sulfate can become fully neutralized.

Response: We don't guess that process of aging is just production of ammonium sulfate without its further modifications. Surely, formation of sulfate through bisulfate is a major way, however, changes are not stopped after formation of sulfate. First, when the ammonium sulfate is in a solution the ions do not bond each other but they are in a form of NH_4^+ and SO_4^{2-} . At the same time, NH_3 from gas phase is absorbed into the droplet. During evaporation of water and part of ammonia, the lighter ammonia is evaporated more and aerosol is enriched by heavier NH_4^+ . It implies, the older the aerosol the more ^{15}N in ammonium sulfate. Second, as shown by recent research (Weber et al., 2016) sulfate is probably not a definitive compound that is not undergo to further changes in time. There probably exist an equilibrium between sulfate and bisulfate which can also affect subsequent changes in gas/particle partitioning of ammonia. Based on this, we added following sentences to related paragraph in subsection 3.3:

"Finally, summer values of $\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratio below 2 indicate that SO_4^{2-} in aerosol particles at high summer temperatures may not be completely saturated with ammonium but it can be composed from mixture of $(\text{NH}_4)_2\text{SO}_4$ and NH_4HSO_4 (Weber et al., 2016). The equilibrium reaction between these two forms of ammonium sulfates related to temperature oscillation during a day and due to vertical mixing of the atmosphere is a probable factor which leads to increased values of $\delta^{15}\text{N}$ in early summer."

Line 436. Oxidation by ozone indeed makes organic matter enriched in heavier carbon, because ozone attacks unsaturated bonds and those involving lighter carbon are preferentially broken releasing "light" CH_3 fragments making the bulk matter heavier.

Response: Thank you for this supportive comment. At the end of sentence we added new following reference related to enrichment of ^{13}C by photochemical processing of aqueous aerosols – see Pavuluri and Kawamura (2016) in references – and we also modified the sentence to following: *“As seen in Table 3, summertime positive correlations of $\delta^{13}\text{C}$ with ozone ($r=0.66$) and temperature (0.39) indicate oxidation processes that can indirectly lead to an enrichment of ^{13}C in organic aerosols that are enriched with oxalic acid (Pavuluri and Kawamura, 2016).”*

Line 444. “...depleted products in summer”. Is this contradictory to the above paragraph?

Response: It seems in contradictory to the above paragraph, however, these are two different things. Even if summer ^{13}C is most depleted compared to other seasons (probably due to different sources) there is a possible indirect oxidation process for their enrichment. Based on correlation analysis, this process is relevant only in summer, however, this enrichment is not strong enough to reach average $\delta^{13}\text{C}$ values during other seasons. The time series in Fig. 1 show the lowest $\delta^{13}\text{C}$ values in a mid of June and slowly increasing enrichment of ^{13}C during rest of summer, which also support this process. We did not make any revisions in MS related to this comment.

Line 454. EC is a minor fraction of TC, so correlations are a bit pointless as EC isotope content contributes little to the TC isotope content.

Response: It is true that EC is a minor fraction of TC, however, in case of our data EC contributes by 19% on average during all seasons, which is not negligible. Interpretations of the results related to EC are supported also by other correlations, namely between $\delta^{13}\text{C}$ and NO_2 , NO_3^- and EC/TC ratio, so we believe that it is not pointless. However, it is possible that these results can be biased by lower content of EC in TC thus we modified part of last sentence at the end of related paragraph as follows: *“This result can be biased by the fact that EC constitutes on average 19% of TC during all seasons. However, it is consistent with positive correlations between $\delta^{13}\text{C}$ and gaseous NO_2 , as well as particulate nitrate, which is also significant in autumn to spring. This result is also supported by the negative correlation between $\delta^{13}\text{C}$ and EC/TC ratio ($r=-0.51$), which is significant only in summer.”*

Line 476. Sulfate cannot be emitted, reword.

Response: Changed to *“formation of sulfates”*

Line 507. Dry deposition of fine particulate matter is negligible making this assertion pure speculation. Dry deposition of ammonia can only occur on surfaces - were these frosted surfaces? Confusing interpretation, please reword.

Response: Thank you for this comment. It is right that dry deposition of fine particulate matter is negligible, however, in mentioned part of text we discussed possible dry deposition of gaseous ammonia. Moreover, given sentence is a reference to observations of Savard et al. (2017). Throughout the Event, the temperature was below 0°C (see Fig.9), so frosted surfaces could be possible on the ground surface, which can support deposition but also decrease fluxes of ammonia from e.g. water surfaces or soil. Also the results indicate that during the Event gradually rose deficit of ammonia and

after some time the main source of ammonia were probably agricultural emissions from farms whose emissions of NH₃ are not as affected by low temperatures. Nevertheless, because we did not measure ammonia fluxes during the campaign, the conclusions on dry deposition heavier isotope of ammonia may sound little bit speculative (as the reviewer mentioned). In this sense we changed text in subsection 3.4 and in Conclusions.

New text in subsection 3.4. (original lines 507-511): *“They interpreted the result as a preferential dry deposition of heavier isotopic ¹⁵NH₃ species during the cold period, whereas lighter ¹⁴NH₃ species preferentially remains in the atmosphere. However, cold weather can also lead to a decline of ammonia fluxes from aerosol water surfaces, soil, etc. (Roelle and Aneja, 2002), which generally result in a deficit of ammonia in the atmosphere. Emissions from farms are not as limited by low temperature and are thus a main source of ammonia in this deficiency state. The removal of NH₃ leads to a non-equilibrium state between the gas and aerosol phases.”*

New text in Conclusions (original lines 577-582): *“The winter Event characterized by ¹⁵N depletion was probably caused by preferential unidirectional reactions between isotopically light ammonia, originated mainly from agriculture emissions, and sulfuric acid, resulting in (NH₄)₂SO₄ and NH₄H₂PO₄. This process was probably supported by long-term cold weather leading to a deficiency of ammonia in the atmosphere (due to dry deposition and/or low fluxes), and subsequent suppression of nitrate to partitioning in aerosol.”*

Table 2&3. Omit insignificant correlation as they are meaningless.

Response: We would like to keep all the values in Tables 2 and 3, however, we removed insignificant correlations from the text - see our responses for major comments in more details.

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Response to anonymous Referee #2 comments

This paper presents seasonal variations of $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ in ambient aerosol collected in Košetice (Central Europe) between 27 September 2013 and 9 August 2014. The authors show an impressive series of measurements aiming to investigate sources and processing of the fine fraction of aerosol at a rural background site. This study using two-isotope analysis is very suitable for this goal.

The use of multiple isotope ratios for the study of atmospheric pollution and the chemistry of organic compounds in the atmosphere is a newly emerging tool. The manuscript contributes to scientific progress within the scope of the journal; therefore, it is suitable to be published for discussions in ACP. Both description and discussion of measurements are well founded. Unfortunately, the presentation is not on the same level, therefore it needs to be substantially improved before publishing.

Response: We would like to thank the reviewer for his valuable and helpful comments. Based on this, we have changed a part of the manuscript which led to an improvement in the final text. Responses to specific comments are below.

General comments:

1) The authors discuss the benefits of using isotopes in the atmospheric research. These can give some hints to information, which is not available from concentration measurements, such as the impact of sources vs. processing on measured delta values. I miss though a discussion on the current limitations of using isotope ratio measurements for the above mentioned purpose. This omission might be the reason why the interpretation sounds sometimes so futile.

Example: Lines 262-263 'In the case of our data, mixing of all of these factors probably had an influence on the nitrate isotopic composition during different parts of the year.'

Reformulate!

Response: Thank you for this comment. We added following text referring to the current limitations of using isotope ratio measurements in first paragraph of Introduction chapter.

"However, studies based on single isotope analysis have their limitations (Meier-Augenstein and Kemp, 2012). Those include an uncertainty when multiple sources or different processes are present, whose measured delta values may overlap (typically in the narrower $\delta^{13}\text{C}$ range). Another factor are isotope fractionation processes which may constrain the accuracy of source identification (Xue et al., 2009). Using isotope analysis on multiple phases (gas and particulate matter) or multiple isotope analysis can overcome these problems and may be useful to constrain the potential sources/processes."

Specific text on lines 262-263 were changed to the following sentence: *"In our study, it is most likely that all these factors contributed, to a certain extent, to the nitrogen isotopic composition of NO_3^- throughout the year."*

2) The introduction should make the reader aware of the importance of using multiple isotopes (literature sources are required), e.g. for constraining potential sources. The sentence on the Lines 85-86 is too late and too less. A proper foreword would bring more structure in the discussion from Lines 59-83. Here the authors must clearly differentiate between single and multiple isotope analyses.

Response: We extended Introduction chapter about examples of studies using multi isotope analyses. The sentence from the lines 85-86 we slightly modified and moved at the end of this paragraph. New text related to studies with multiple isotope measurements in Introduction chapter is following:

“Recently, the multiple isotope approach was applied in several studies by using $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ measurements. Specifically, the $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ composition of aerosol (along with other supporting data) was used to identify the sources and processes on marine sites in Asia (Bikkina et al., 2016; Kunwar et al., 2016; Miyazaki et al., 2011; Xiao et al., 2018). Same isotopes were used to determine the contribution of biomass burning to organic aerosols in India (Boreddy et al., 2018) and in Tanzania (Mkoma et al., 2014), or to unravel the sources of aerosol contamination at Cuban rural and urban coastal sites (Morera-Gómez et al., 2018). These studies show the potential advantages of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ isotope ratios to characterize aerosol types and to reveal the underlying chemical processes that take place in them.”

We also added data on other isotope analyzes (if were performed) to distinguish single- and multi-isotope studies in paragraph related to European studies

3) Separate Spearman from Pearson correlation coefficients. For that purpose, label them for each use (e.g. in Line203).

Response: Thank you for this comment. We identified in text few Pearson's correlation coefficients (connected with Figures 2, 3 and 7) instead of Spearman's ones. Although each of these coefficients provides different information (Pearson benchmarks linear relationship, Spearman benchmarks monotonic relationship), the values in our work are same or similar (e.g. for TC vs.TN is $r(P)$: 0.70 and $r(S)$: 0.71). Based on this, we decided to use only Spearman correlation coefficients in this work. Changes were made in Figures 2, 3 and 7, and related text (original lines 203, 213). Currently, Spearman's correlations are used throughout the document so there is no need to differentiate it from Pearson's correlations.

4) Name the described variables throughout the manuscript!

Some examples: Line122: Replace 'Determination of TC, TN and their stable isotopes'

by 'Determination of TC, TN concentrations and their stable isotope ratios'

Line123: Replace 'For the TC and TN analyses' by ' For the TC and TN concentration

and isotopic ratio measurements'

Response: For a text clarification, there were changed variables description on following lines: 122, 123, 202, 205, 290, 545,

5) Vague statements should be replaced by precise explanations throughout the paper.

An example: Line382: specify the 'secondary processes'

Response: We are sorry for vague statements. We rephrased the text as below.

Statement on line 382 was based on work of Widory (2007). We changed the previous sentence to following: *“Similarly, the contradictory dependence between $\delta^{15}\text{N}$ and TN in summer and winter was observed by Widory (2007) in PM10 samples from Paris. Widory (2007) connected this result with different primary nitrogen origin (road-traffic emissions in summer and no specific source in winter) and following secondary processes associated with isotope fractionation during degradation of atmospheric NOx leading to two distinct pathways for ^{15}N enrichment (summer) and depletion (winter).”*

R: 6) Generally: swap the negative numbers in ranges. The lower numbers stay first.

Examples: Line520 -40 to -28permil and Line522 -38 to -22permil

Response: Ranges of negative numbers were swapped on original lines 338, 520, 522 and 533. Thank you for your notice.

Specific comments:

Lines54-57: Reformulate! The OC/EC ratios are very different in aerosol, depending on its sources. Moreover, make more sentences of this single one. Differentiate between equilibrium and kinetic isotopic effect. Guide the reader through that by giving some information on corresponding fractionation (non-equilibrium partitioning causes much lower fractionation than chemical reactions. Contrarily, equilibrium fractionation might be significant).

Response: We changed the Introduction chapter with the text related to isotopes in carbonaceous aerosols and we also inserted a new paragraph on isotope fractionation:

New text related to carbonaceous aerosols:

“Total carbon in aerosol is usually divided into elemental carbon (EC) and organic carbon (OC), where OC forms the major part of TC (e.g., Mbengue et al., 2018). Although EC is more or less inert to chemical changes, slightly different $\delta^{13}\text{C}$ in EC originating from primary emissions are described (Kawashima and Haneishi, 2012). OC represents a wide variety of organic compounds which can originate from different sources with different ^{13}C content resulting in different $\delta^{13}\text{C}$ values in bulk of emissions. Changes in isotopic ratio of $\delta^{13}\text{C}$ in OC (and thus also TC) can subsequently affect chemical reactions where isotope fractionations via the kinetic isotope effect (KIE) usually dominate the partitioning between gas and aerosol (liquid/solid) phases (e.g. Zhang et al., 2016).”

New paragraph related to isotope fractionation:

“Isotopes are furthermore altered mainly by kinetic and/or equilibrium fractionation processes. Kinetic isotope effects (KIE) occur mainly during unidirectional (irreversible) reactions but also diffusion or during reversible reactions that are not yet at equilibrium (Gensch et al., 2014). Owing to KIE, reaction products (both gasses and particles) are depleted in the heavy isotope relatively to the reactants, and this effect is generally observed in organic compounds (Irei et al., 2006). If the partitioning between phases is caused by non-equilibrium processes (such as e.g. absorption), the isotopic fractionation is small and lower than that caused by chemical reactions (Rahn and Eiler, 2001). Equilibrium isotope effects occur in reversible chemical reactions or phase changes if the system is in equilibrium. Under such conditions, the heavier isotope is bound into the compounds where the total energy of the system

is minimized and the most stable. Equilibrium effects are typical for inorganic species and usually temperature dependent.”

Line87: No need to introduce TC and TN. It happened already in Lines12-13

Response: Edited and only shortcuts were kept.

Line127: I don't understand. Is the oven temperature 1000 °C? How can the marble burn, if that needs 1400 °C?

Response: Theory is that burning tin should locally increase temperature around the sample to approximately 1400°C, however, this is not so important and it can be also confusing so we deleted temperature 1400°C from the MS.

Line131: What does 'parts' means? Give the approximate fraction in %.

Response: At this point, there is an auto-dilution system on the ConFlo IV interface, which is applied to each gas species matching sample and reference gas intensities. This dilution is automatic and the device dynamically reacts to the sample volume. Therefore, it is not possible to specify the exact part of the sample. For this reason, we decided to shorten the sentence to the following form:

“~~Parts of~~ CO₂ and N₂ were then transferred into an isotope ratio mass spectrometer (IRMS; Delta V, Thermo Fisher Scientific) through a ConFlo IV interface to monitor ¹⁵N/¹⁴N and ¹³C/¹²C ratios.”

Lines135-139: Mention that the final delta values are expressed relatively to the international standards and not to the 'working' standard.

Response: That's a good point. Sentence before equations was extended to following form:

'Subsequently, $\delta^{15}\text{N}$ of TN and $\delta^{13}\text{C}$ of TC were calculated using the following equations and the final δ values are expressed in relation to the international standards:'

Line146: The loads on the quartz filter are meant here of course.

Response: Yes, you are right that the loads on quartz filters was analyzed. The sentence was changed in this sense.

Lines198-200: Move these sentences to the first paragraph, they don't belong to Fig.1.

Response: You are right, these sentences belong to Table 1 so we just changed link to Fig.1 to Tab.1 at the end of this paragraph.

Lines218-219: Reformulate: 'but they are still in line with the linear fitting of all annual data'. This is not appropriate.

Response: The sentence was reworded to the following: *“The winter Event measurements show the highest $\delta^{13}\text{C}$ and the lowest $\delta^{15}\text{N}$, but a linear fit does not show a significant differences as compared to rest of the data (Fig. 2, right).”*

Lines290-291: Reformulate! Either state that the samples containing the highest NO_3^- concentration show a $\delta^{15}\text{N}$ of..., or fit a histogram plot showing a peak of measurements with NO_3^- concentrations higher than... at a delta value of 14 ± 1 permil.

Response: Thank you for this comment. You are right that statement on lines 290-291 *“The $\delta^{15}\text{N}$ shows a peak at approximately $14\pm 1\%$...”* is not exact, and is the result of estimation based on exponential curves in Figure 4. So newly, we took samples with NO_3^- concentrations higher than $6\ \mu\text{g}/\text{m}^3$ ($n=5$) and we calculated an average $\delta^{15}\text{N}$ value from these samples. It results in new value of $\delta^{15}\text{N}$ ($13.3\pm 0.7\%$), we used this calculated value instead of $14\pm 1\%$ in whole text.

New text on lines 290-291 is following: *“Samples with the highest NO_3^- concentrations ($>6\ \mu\text{g}/\text{m}^3$, $n=5$) show an average $\delta^{15}\text{N}$ of $13.3\pm 0.7\%$.”*

Lines300-307: The paragraph should be moved upward to Fig. 3.

Response: The paragraph on lines 300-307 relates to the previous one, where the results in Figure 4 are commented. For this reason, we would like keep this paragraph in the current position.

Lines338-349: Completely rearrange! Suggestion: start with a statement ‘The measured TC $\delta^{13}\text{C}$ ranged between.... These values are ... (in which part?) situated in the reported ranges... (here give an overall range. for that take the information from e.g. the review by Gensch et al. 2014). This broad range can be explained by... (plants, marine, combustion sources... whatever). (At this point bring the similarity to other european reported values).’

Response: Thank you for this suggestion. Based on this, paragraph was rearranged as below:

“The $\delta^{13}\text{C}$ of TC ranged from -28.9 to -25.4% (Fig. 6) and the lowest $\delta^{13}\text{C}$ we observed in field blank samples (mean -29.2% , $n=7$), indicating that the lowest summer values in particulate matter were close to gas phase values. Our $\delta^{13}\text{C}$ values are within the range reported for particulate TC (-29% to -15%) as summarized by Gensch et al. (2014). The lowest values are associated with fine particles after combustion and transport (Ancelet et al., 2011; Widory, 2006) while the highest values are associated with the coarse fraction and carbonate contribution (Kawamura et al., 2004). This broad range can be explained by the influence of marine aerosols (Ceburnis et al., 2016), different anthropogenic sources (e.g., Widory et al., 2004), as well as different distributions of C3 and C4 plants (Martinelli et al., 2002) resulting in different $\delta^{13}\text{C}$ values in the northern and southern hemispheres (Cachier, 1989). The $\delta^{13}\text{C}$ values at the Košetice site fall within the range common to other European sites. For example, a rural background site in Vavihill (southern Sweden, range -26.7 to -25.6% , Martinsson et al. (2017)), urban Wroclaw (Poland, range -27.6 to -25.3% , Górka et al. (2014)), different sites (urban, coastal, forest) in Lithuania (East Europe, Masalaite et al., 2015, 2017), as well as urban Zurich (Switzerland, Fisseha et al. (2009)).”

Line349: Replace 'The d13C values are significantly smaller than those of d15N due to' by 'The range of TC d13C values is significantly smaller than that of TN d15N due to'

Response: The sentence was changed based on the comment as bellow: "The range of TC $\delta^{13}C$ values is significantly narrower than that of TN $\delta^{15}N$ due to..."

Lines358-359: This comparison is confusing: what do you mean? Similar to what? Do you refer the first or the second sentence?

Response: The comparison was between first and third sentence. Second sentence was moved to first paragraph of section 3.2, so now it should be clearer. See first paragraph of subsection 3.2 in the revised MS.

Lines365-370: Change the order of these two sentences. Describe first the observations and then give the explanation.

Response: The order of sentences was changed. See end paragraph of subsection 3.2 in the revised MS.

Line 375: Replace 'these isotopes' with 'isotope distributions'.

Response: Replaced

Lines379-380: Not the changes in aerosol chemistry are different, but the chemistry itself.

Response: Word "changes" was deleted.

Lines386-391: Change the order of the first two sentences. The third one describes the first not the second one.

Response: The order of sentences was changed.

Lines415-422: Lack of clarity! Reformulate, by bringing some structure in it: starting at high NH₄/SO₄ down to 2 and lower than 2! For each range: particle components, processes (e.g. NH₃ deficit in gas phase at ratios <2), seasonal dependence.

Response: Thank you for your suggestion. Based on this comment, we decided to completely change this paragraph to the following:

"Figure 8 shows an enrichment of ^{15}N as a function of the molar ratio of NH_4^+/SO_4^{2-} . The highest NH_4^+/SO_4^{2-} ratios, showing an ammonia rich atmosphere, were observed during winter, late autumn and early spring along with high abundance of NO_3^- that is related to favorable thermodynamic conditions during heating season and enough ammonia in the atmosphere. Gradual decreasing molar ratios of NH_4^+/SO_4^{2-} during spring indicate a gradual increase of ambient temperatures and therefore worsened thermodynamic conditions for NO_3^- formation in aerosol phase, which was accompanied by

a visible decrease in the nitrate content in aerosols (Fig. 8). The increase of temperatures finally leads to the $\text{NH}_4^+/\text{SO}_4^{2-}$ ratio reaching 2 at the turn of spring and summer. Finally, summer values of $\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratio below 2 indicate that SO_4^{2-} in aerosol particles at high summer temperatures may not be completely saturated with ammonium but it can be composed from mixture of $(\text{NH}_4)_2\text{SO}_4$ and NH_4HSO_4 (Weber et al., 2016). The equilibrium reaction between these two forms of ammonium sulfates related to temperature oscillation during a day and due to vertical mixing of the atmosphere is a probable factor which leads to increased values of $\delta^{15}\text{N}$ in early summer. Ammonia measurements, that were carried out at the Košetice site until 2001, showed that NH_3 concentrations in summer were slightly higher than in winter (http://portal.chmi.cz/files/portal/docs/uoco/isko/tab_roc/2000_enh/CZE/kap_18/kap_18_026.html), which supports temperature as a main factor influencing $\text{NH}_4^+/\text{SO}_4^{2-}$ ratio at Košetice. In this context, we noticed that 25 out of 33 summer samples have molar ratios of $\text{NH}_4^+/\text{SO}_4^{2-}$ below 2, and the remaining samples are approximately 2, and the relative abundance of NO_3^- in PM1 in those samples is very low (ca. 1.7 %)."

Lines429-434: Too abrupt! Start with the observation of similar gaseous NH_3 in summer and winter. Describe what a thermodynamic equilibrium would mean for the particles and how would this be reflected in the delta values. Measurements show a different situation -> more organics in summer...

Response: Thank you for this notice. We moved the sentence, about similar gaseous NH_3 concentrations in summer and winter at the Košetice site, to paragraph above (see response to previous comment). The lines have been reformulated into the following form:

„ In thermodynamic equilibrium, partitioning between gas (NH_3) and aerosol (NH_4^+) phases should result in even larger $\delta^{15}\text{N}$ values of particles in summer, however, measurements show a different situation. Summer $\delta^{15}\text{N}$ values are highest but further enrichment was stopped. Moreover, we observed a positive (and significant) correlation between temperature and $\delta^{13}\text{C}$ ($r=0.39$) only in summer, whereas the correlation coefficient of $\delta^{15}\text{N}$ vs. temperature is statistically insignificant, suggesting that while values of $\delta^{15}\text{N}$ reached their maxima, the $\delta^{13}\text{C}$ can still grow with even higher temperatures due to the influence of organics in summer season.“

Lines482-484: Very confuse sentence. Reformulate!

Response: The sentence is reworded to the following: *“During the Event, $\delta^{15}\text{N}$ correlates positively with NO_3^- ($r=0.96$) and NO_3^- -N/TN (0.98). Before the Event, we also observed the highest values of $\delta^{15}\text{N}$ at approximately 13.3‰, which we previously interpreted by the emissions from domestic heating via coal and/or biomass burning.“*

Lines570-574: The winter observation should stay before the summer ones. In that way, the flow is more coherent (e.g. no need to explain lower values of TN d15N when there are high fraction of nitrates.).

Response: In summary and conclusions, we wanted to discuss all seasons for the first time. After summary related to seasonal data follow conclusions related to winter Event. This is reason why the winter data are discussed after summer data. It seems more logic to us if the winter Event summary

follows winter data than to discuss first about winter, then continue about summer season, and after that return to discuss about the winter Event. So we prefer not to move this paragraph.

Editorial revisions:

The used English is not optimal. I do not give any editorial advises! My only suggestion is that this manuscript MUST be carefully revised by a native speaker. The work is too good to risk to make the reader hostile due to the language.

Response: We are sorry for inconvenience with English. In fact, the text was checked by the professional language service before the first submission so we expected it should be alright. As the final step after this review process, we sent again the manuscript for English corrections to Sean Mark Miller who is a native speaker and professional corrector.

The manuscript is 'peppered' with:

1) Wrong prepositions

- Lines43-44 'Key processes in the atmosphere, which are involved WITH climate changes, air quality, rain events (Fuzzi et al., 2015) or visibility (Hyslop, 2009), are strongly influenced by aerosols.' –

Lines391-392 ' Although Savard et al. (2017) reported a similar negative $\delta^{15}\text{N}$ in NH_4^+ dependence AT temperatures in Alberta (Canada),...' Also the word order is wrong.

Response: Text with above mentioned prepositions was changed.

Lines43-44: *"Aerosols have a strong impact on key processes in the atmosphere associated with climate change..."*

Lines391-392: *"Although Savard et al. (2017) reported a similar negative temperature dependence for $\delta^{15}\text{N}$ in NH_4^+ in Alberta (Canada), ..."*

2) Unhandy expressions

- Lines325-328:' During the domestic heating season with the highest concentrations of NO_3 and NH_4^+ , we can observe a significant increase in OrgN with $\delta^{15}\text{N}$ again at approximately 14‰ which implies that the isotopic composition of OrgN is determined by the same process during maximal NO_3 -concentrations, that is, emissions from domestic heating.'

Response: Sentence on lines325-328 was shortened to: *"During the domestic heating season with the highest concentrations of NO_3^- and NH_4^+ , we can observe a significant increase in OrgN with $\delta^{15}\text{N}$ again at approximately 13.3‰, which implies that the isotopic composition of OrgN is determined by the same source."*

3) Long, confusing sentences

Lines361-365 or Lines391-396. In these cases it helps to divide into more clear sentences.

Response: We divided above mentioned long sentences and also others in the revised MS.

Newly for line 361-365: “Similarly, at the Košetice station, different carbonaceous aerosols were observed during the heating season (Oct.–Apr.) than in summer (Mbengue et al., 2018; Vodička et al., 2015). Moreover, winter aerosols at the Košetice site were probably affected by not only biomass burning but also coal burning (Schwarz et al., 2016), which can result in higher carbon contents and more ^{13}C enriched particles (Widory, 2006).”

Newly for lines 391-396: “Although Savard et al. (2017) reported a similar negative temperature dependence for $\delta^{15}\text{N}$ of NH_4^+ in Alberta (Canada), most studies reported a positive temperature dependence for $\delta^{15}\text{N}$ of NH_4^+ that is stronger than that for $\delta^{15}\text{N}$ of NO_3^- (e.g., Kawashima and Kurahashi, 2011; Kundu et al., 2010). The reason is that NH_3 gas concentrations are higher during warmer conditions, and thus the isotopic equilibrium exchange reaction, i.e., $\text{NH}_3(\text{g}) \leftrightarrow \text{NH}_4^+(\text{p})$, which leads to ^{15}N enrichment in particles, is more intensive.”

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Seasonal study of stable carbon and nitrogen isotopic composition in fine aerosols at a Central European rural background station

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Abstract. ~~A study~~ ~~Determinations~~ of ~~the~~ stable carbon isotope ratios ($\delta^{13}\text{C}$) of total carbon (TC) and ~~the~~ nitrogen isotope ratios ($\delta^{15}\text{N}$) of total nitrogen (TN) were carried out for fine aerosol particles (PM1) collected ~~every two days withon a 24 h sampling period~~ ~~daily basis~~ at a rural background site in Košetice (Central Europe) ~~from between 27 September 27, 2013, to and 9 August 9, 2014~~ (n=146). We found a seasonal pattern for both $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$. The seasonal variation in $\delta^{15}\text{N}$ was ~~characterized by lower~~ ~~more pronounced, with ^{15}N depleted~~ values (av. $13.1\pm 4.5\%$) in winter and ~~higher ^{15}N enriched~~ values ($25.0\pm 1.6\%$) in summer. Autumn and spring ~~were~~ ~~are~~ transition periods when the isotopic composition gradually changed due to ~~the changing~~ ~~different~~ sources and the ambient temperature. The seasonal variation in $\delta^{13}\text{C}$ was less pronounced but more depleted in ^{13}C in summer ($-27.8\pm 0.4\%$) ~~as~~ compared to winter ($-26.7\pm 0.5\%$).

~~A~~ ~~Major controls of the seasonal dependencies were found based on a~~ comparative analysis with water-soluble ions, organic carbon, elemental carbon, trace gases and meteorological parameters (mainly ambient temperature) ~~has shown major associations with the isotopic compositions, which enlightened the affecting processes.~~ A comparison of $\delta^{15}\text{N}$ with NO_3^- , NH_4^+ and organic nitrogen (OrgN) revealed that although a higher content of NO_3^- was associated with a decrease in ~~the~~ $\delta^{15}\text{N}$ ~~of values in~~ TN, NH_4^+ and OrgN ~~caused increases~~ ~~had the opposite influences~~. The highest concentrations of nitrate, mainly represented by NH_4NO_3 , ~~related to~~ ~~originated from~~ the emissions from biomass burning, leading to ~~an average lower~~ $\delta^{15}\text{N}$ ~~of TN (13.3%)~~ ~~values of approximately 14%~~ in winter. During spring, the percentage of NO_3^- in PM1 decreased. ~~An,~~ ~~and~~ ^{15}N enrichment ~~of ^{15}N~~ was probably driven by ~~the~~ equilibrium exchange between the gas and aerosol phases ($\text{NH}_3(\text{g}) \leftrightarrow \text{NH}_4^+(\text{p})$), ~~which is~~ ~~as~~ supported by the increased ambient temperature. This equilibrium was suppressed in early summer when the ~~$\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratios~~ ~~of $\text{NH}_4^+/\text{SO}_4^{2-}$~~ reached 2, and ~~the~~ nitrate partitioning in aerosol was negligible ~~due~~. ~~During summer, kinetic reactions probably were the primary processes as opposed to the increased ambient temperature. Summertime gas-aerosol equilibrium on a nitrogen level. However, summertime~~ $\delta^{15}\text{N}$ values were ~~among some of~~ the highest ~~observed, probably~~ suggesting the aging of ammonium

37 sulfate and OrgN aerosols. Such aged aerosols can be coated by organics in which ^{13}C enrichment takes
38 place by the photooxidation process. This result was supported by ~~athe~~ positive correlation of $\delta^{13}\text{C}$ with
39 ambient temperature and ozone, as observed in the summer season.

40 During winter, we observed an event with the lowest $\delta^{15}\text{N}$ and highest $\delta^{13}\text{C}$ values. The winter *Event*
41 ~~occurred in was connected with~~ prevailing southeast air masseswinds. Although the higher $\delta^{13}\text{C}$ values
42 probably originated from biomass burning particles, the lowest $\delta^{15}\text{N}$ values were probably associated
43 with agriculture emissions of NH_3 under low temperature conditions (~~<that were below 0°C~~).

45 1. Introduction

46
47 ~~Aerosols have a strong impact on key~~Key processes in the atmosphere ~~associated, which are involved~~
48 with climate ~~change~~changes, air quality, rain ~~patterns and visibility~~events (Fuzzi et al., 2015; Hyslop,
49 2009).~~(Fuzzi et al., 2015) or visibility (Hyslop, 2009), are strongly influenced by aerosols.~~ Because
50 these processes are still insufficiently understood, they are studied intensively. One approach to explore
51 chemical processes taking place in atmospheric aerosols is the application of stable carbon ($\delta^{13}\text{C}$) and
52 nitrogen ($\delta^{15}\text{N}$) isotope ratios. These isotopes can provide unique ~~insight~~information on source
53 emissions ~~along together~~ with physical and chemical processes in the atmosphere (Gensch et al., 2014;
54 Kawamura et al., 2004), as well as atmospheric composition in history (Dean et al., 2014). However,
55 studies based on single isotope analysis have their limitations (Meier-Augenstein and Kemp, 2012).
56 Those include an uncertainty when multiple sources or different processes are present, whose measured
57 delta values may overlap (typically in the narrower $\delta^{13}\text{C}$ range). Another factor are isotope fractionation
58 processes which may constrain the accuracy of source identification (Xue et al., 2009). Using isotope
59 analysis on multiple phases (gas and particulate matter) or multiple isotope analysis can overcome these
60 problems and may be useful to constrain the potential sources/processes.

61 Generally, isotopic~~isotopic~~ composition is affected by both primary emissions (e.g., Heaton, 1990;
62 Widory, 2006) and secondary processes (e.g., Fisseha et al., 2009b; Walters et al., 2015a). Isotopes are
63 furthermore altered mainly by kinetic and/or equilibrium fractionation processes. Kinetic isotope effects
64 (KIE) occur mainly during unidirectional (irreversible) reactions but also diffusion or during reversible
65 reactions that are not yet at equilibrium~~Both $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ values are influenced by kinetic and~~
66 ~~equilibrium isotope fractionation that takes place in the atmosphere. In the case of nitrogen, ^{15}N is~~
67 ~~generally depleted in gas phase precursors (ammonia, nitrogen oxides) but is more enriched in ions~~
68 ~~(NH_4^+ , NO_3^-) in rainfall and most enriched in particulate matter and dry deposition (Heaton et al., 1997).~~
69 In the case of carbon, the major form is organic carbon (OC), which is composed of large numbers of
70 organic compounds where isotope fractionations via the kinetic isotope effect (KIE) usually dominating
71 the partitioning between gas and aerosol (liquid/solid) phases (Gensch et al., 2014). Owing to KIE,
72 reaction products (both gasses and particles) are depleted in the heavy isotope relatively to the reactants,

73 and this effect is generally observed in organic compounds (Irei et al., 2006). If the partitioning between
74 phases is caused by non-equilibrium processes (such as e.g. absorption), the isotopic fractionation is
75 small and lower than that caused by chemical reactions (Rahn and Eiler, 2001). Equilibrium isotope
76 effects occur in reversible chemical reactions or phase changes if the system is in equilibrium. Under
77 such conditions, the heavier isotope is bound into the compounds where the total energy of the system
78 is minimized and the most stable. Equilibrium effects are typical for inorganic species and usually
79 temperature dependent.

80 Regarding to the isotopic distribution in individual phases, ^{15}N is generally depleted in gas phase
81 precursors (ammonia, nitrogen oxides) but is more enriched in ions (NH_4^+ , NO_3^-) in rainfall and the
82 most enriched in particulate matter and dry deposition (Heaton et al., 1997; Ti et al., 2018). Total
83 nitrogen usually consists of the three main components, NO_3^- , NH_4^+ and/or organic nitrogen (OrgN),
84 and thus, the final $\delta^{15}\text{N}$ value in TN can be formulated by the following equation:

$$\delta^{15}\text{N}_{\text{TN}} = \delta^{15}\text{N}_{\text{NO}_3^-} * f_{\text{NO}_3^-} + \delta^{15}\text{N}_{\text{NH}_4^+} * f_{\text{NH}_4^+} + \delta^{15}\text{N}_{\text{OrgN}} * f_{\text{OrgN}}$$

86 where $f_{\text{NO}_3^-} + f_{\text{NH}_4^+} + f_{\text{OrgN}} = 1$ and f represents the fractions of nitrogen from NO_3^- , NH_4^+ and OrgN in
87 TN, respectively.

88 Total carbon in aerosol is usually divided into elemental carbon (EC) and organic carbon (OC), where
89 OC forms the major part of TC (e.g., Mbengue et al., 2018). Although EC is more or less inert to
90 chemical changes, slightly different $\delta^{13}\text{C}$ in EC originating from primary emissions are described
91 (Kawashima and Haneishi, 2012). OC represents a wide variety of organic compounds which can
92 originate from different sources with different ^{13}C content resulting in different $\delta^{13}\text{C}$ values in bulk of
93 emissions. Changes in isotopic ratio of $\delta^{13}\text{C}$ in OC (and thus also TC) can subsequently affect chemical
94 reactions where isotope fractionations via the kinetic isotope effect (KIE) usually dominate the
95 partitioning between gas and aerosol (liquid/solid) phases (e.g. Zhang et al., 2016).

96
97 Many studies have been conducted on $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ in particulate matter (PM) in Asia (e.g., Kundu et
98 al., 2010; Pavuluri et al., 2015b; Pavuluri and Kawamura, 2017) and ~~the Americas~~ America (e.g.,
99 Martinelli et al., 2002; Savard et al., 2017). Recently, the multiple isotope approach was applied in
100 several studies by using $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ measurements. Specifically, the $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ composition of
101 aerosol (along with other supporting data) was used to identify the sources and processes on marine
102 sites in Asia (Bikkina et al., 2016; Kunwar et al., 2016; Miyazaki et al., 2011; Xiao et al., 2018). Same
103 isotopes were used to determine the contribution of biomass burning to organic aerosols in India
104 (Boreddy et al., 2018) and in Tanzania (Mkoma et al., 2014), or to unravel the sources of aerosol
105 contamination at Cuban rural and urban coastal sites (Morera-Gómez et al., 2018). These studies show
106 the potential advantages of $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ isotope ratios to characterize aerosol types and to reveal the
107 underlying chemical processes that take place in them.

108 ~~Only~~ However, only few studies on $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ isotope ratios have been performed in Europe, which
109 are moreover often based on single. ~~European~~ isotope studies on aerosols mainly involve the analysis.

110 ~~Regarding the isotopes of nitrogen, $\delta^{15}\text{N}$ in NO_3^- and/or NH_4^+ .~~ Widory (2007) published a broad study
111 on $\delta^{15}\text{N}$ in TN in PM10 samples from Paris, focusing on ~~seasonality seasons~~ (winter vs. summer) with
112 some specific sources. Freyer (1991) reported the seasonal variation in ~~the~~ $\delta^{15}\text{N}$ of nitrate in aerosols
113 and rainwater as well as gaseous HNO_3 at a moderately polluted urban area in Jülich (Germany).
114 Yeatman et al. (2001a, 2001b) conducted analyses of $\delta^{15}\text{N}$ in NO_3^- and NH_4^+ at two coastal sites from
115 Weybourne, England, and Mace Head, Ireland, focusing on the effects of ~~the~~ possible sources and
116 aerosol size segregation on their formation processes and isotopic enrichment. More recently, Ciężka
117 et al. (2016) reported one-year observations of $\delta^{15}\text{N}$ in NH_4^+ and ions in precipitation at an urban site
118 in Wrocław, Poland, whereas Beyn et al. (2015) reported seasonal changes in $\delta^{15}\text{N}$ in NO_3^- in wet and
119 dry deposition at a coastal and an urban site in Germany to evaluate ~~the~~ nitrogen pollution levels.
120 Studies on $\delta^{13}\text{C}$ at European sites have been focused more on urban aerosols. Fisseha et al. (2009) used
121 stable carbon isotopes ~~of the different carbonaceous aerosol fractions (TC, black carbon, and water~~
122 ~~soluble and insoluble OC)~~ to determine the sources of urban ~~carbonaceous~~ aerosols in Zurich,
123 Switzerland, during winter and summer. Similarly, Widory et al. (2004) used $\delta^{13}\text{C}$ of TC, ~~along together~~
124 with an analysis of lead isotopes, to study the origin of aerosol particles in Paris (France). Górká et al.
125 (2014) used $\delta^{13}\text{C}$ in TC ~~in conjunction together~~ with PAH analyses for the determination of ~~the~~ sources
126 of PM10 organic matter in Wrocław, Poland, during vegetative and heating seasons. Masalaite et al.
127 (2015) used an analysis of $\delta^{13}\text{C}$ in TC on size-segregated urban aerosols to elucidate carbonaceous PM
128 sources in Vilnius, Lithuania. Fewer studies have been conducted on $\delta^{13}\text{C}$ in aerosols in rural and remote
129 areas of Europe. In the 1990s, Pichlmayer et al. (1998) conducted ~~a multiple~~ isotope analysis ~~of $\delta^{13}\text{C}$~~
130 ~~in OC, $\delta^{15}\text{N}$ in NO_3^- and $\delta^{34}\text{S}$ in SO_4^{2-}~~ in snow and air samples for the characterization of pollutants at
131 high-alpine sites in Central Europe. Recently, Martinsson et al. (2017) published seasonal observations
132 of $\delta^{13}\text{C}$ in TC, ~~along with the $^{14}\text{C}/^{12}\text{C}$ isotope ratio~~ of PM10 at a rural background station in Vavíhill in
133 southern Sweden based on 25 weekly samples.

134
135 ~~These $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ studies show the potential of these isotopes to characterize aerosol types and the~~
136 ~~chemical processes that take place in them.~~ To broaden ~~the multiple~~ ~~this~~ isotope approach over the
137 European continent, we present seasonal variations in $\delta^{13}\text{C}$ of ~~total carbon (TC)~~ and $\delta^{15}\text{N}$ of ~~total~~
138 ~~nitrogen (TN)~~ in the PM1 fraction of atmospheric aerosols at a rural background site in Central Europe.
139 To the best of our knowledge, this is the first seasonal study of these isotopes in this ~~region~~ ~~location~~,
140 and it is one of the most comprehensive isotope studies of a fine fraction of aerosols.

141

142 2. Materials and methods

143 2.1. Measurement site

144

145 The Košetice observatory is ~~a key station~~~~the specialized workplace~~ of the Czech Hydrometeorological
146 Institute (CHMI), ~~focusing~~~~which is focused~~ on air quality and environmental monitoring ~~the quality of~~
147 ~~the environment~~ (Váňa and Dvorská, 2014). The site is located in the Czech Highlands (49°34'24.13"
148 N, 15°4'49.67" E, 534 m ASL) and is surrounded by an agricultural landscape and forests, out of range
149 of major sources of pollution with ~~a very low~~ ~~frequency of traffic~~ density. The observatory is officially
150 classified as a Central European rural background site, which is part of the EMEP, ACTRIS, and GAW
151 networks. A characterization of the station in terms of the chemical composition of fine aerosols during
152 different seasons and air masses is presented by Schwarz et al. (2016) and longtime trends by Mbengue
153 et al. (2018) and Pokorná et al. (2018). As part of a monitoring network operated by the CHMI, the site
154 is equipped with an automated monitoring system that provides meteorological data (wind speed and
155 direction, relative humidity, temperature, pressure, and solar radiation) and the concentrations of
156 gaseous pollutants (SO₂, CO, NO, NO₂, NO_x and O₃).

158 2.2. Sampling and weighing

159
160 Aerosol samples (~~n = 146~~) were collected ~~for 24 h~~ every two days for 24 h from September 27, 2013,
161 to August 9, 2014, using a Leckel sequential sampler SEQ47/50 equipped with a PM1 sampling inlet.
162 Some temporal gaps ~~in sampling~~ were caused by ~~sampler outages and~~ maintenance or power outages
163 resulting in 146 samples during ~~to the almost year-long study~~ ~~sampler~~. The sampler was loaded with pre-
164 baked (3 h, 800°C) quartz fiber filters (Tissuequartz, Pall, 47 mm), and operated at a the flow rate of 2.3
165 m³/h ~~was used~~. In addition, field blanks (n = 7) were also taken for an analysis of the contribution of
166 absorbable organic vapors.

167
168 The ~~mass of~~ PM1 was measured gravimetrically ~~(by gravimetric analysis of each quartz filter before~~
169 ~~and after the sampling)~~ with a microbalance that had ±1 µg sensitivity (Sartorius M5P, Sartorius AG,
170 Göttingen, Germany) in a controlled environment ~~(-)~~. ~~The weighing of samples was performed at~~
171 ~~20±1 °C and 50±3 % relative humidity after filter equilibration for 24 h)~~.

173 2.3. Determination of TC, TN concentrations and their stable isotopes

174
175 For the measurements of total carbon (TC) and nitrogen (TN) and their stable isotope ratios ~~analyses~~,
176 small filter discs (area 0.5 cm², 1.13 cm² or 2.01 cm²) were placed in a pre-cleaned tin cup, shaped into
177 a small marble using a pair of tweezers, and introduced into the elemental analyzer (EA; Flash 2000,
178 Thermo Fisher Scientific) using an autosampler. Inside the EA, samples were first oxidized in a quartz
179 column heated at 1000°C, in which the tin marble burns ~~(~1400°C)~~ and oxidizes all the carbon and
180 nitrogen species to CO₂ and nitrogen oxides, respectively. In the second quartz column, heated to 750°C,

181 nitrogen oxides were reduced to N₂. Evolved CO₂ and N₂ were subsequently separated on a gas
182 chromatographic column, which was installed in EA, and measured with a thermal conductivity detector
183 for TC and TN. ~~Parts of~~ CO₂ and N₂ were then transferred into an isotope ratio mass spectrometer
184 (IRMS; Delta V, Thermo Fisher Scientific) through a ConFlo IV interface to monitor the ¹⁵N/¹⁴N and
185 ¹³C/¹²C ratios.

186

187 An acetanilide external standard (from Thermo ~~Electron Corp~~ ~~electron corp.~~) was used to determine
188 the calibration curves before every set of measurements for ~~calculating the calculation of the right~~
189 ~~values of~~ TC, TN and their ~~isotope values~~ ~~isotopic ratios~~. The δ¹⁵N and δ¹³C values of the acetanilide
190 standard were 11.89‰ (relative to the atmospheric nitrogen) and -27.26‰ (relative to Vienna Pee Dee
191 Belemnite standard), respectively. Subsequently, the δ¹⁵N of TN and δ¹³C of TC were calculated using
192 the following equations ~~and the final δ values are expressed in relation to the international standards:~~

193

$$194 \delta^{15}\text{N} (\text{‰}) = [({}^{15}\text{N}/{}^{14}\text{N})_{\text{sample}} / ({}^{15}\text{N}/{}^{14}\text{N})_{\text{standard}} - 1] * 1000$$

$$195 \delta^{13}\text{C} (\text{‰}) = [({}^{13}\text{C}/{}^{12}\text{C})_{\text{sample}} / ({}^{13}\text{C}/{}^{12}\text{C})_{\text{standard}} - 1] * 1000$$

196

197 2.4. Ion chromatography

198

199 ~~The loads on the quartz~~ ~~Quartz~~ filters ~~was~~ ~~were~~ further analyzed by using a Dionex ICS-5000 (Thermo
200 Scientific, USA) ion chromatograph (IC). The samples were extracted using ultrapure water with
201 conductivity below 0.08 μS/m (Ultrapur, Watrex Ltd., Czech Rep.) for 0.5 h using an ultrasonic bath
202 and 1 h using a shaker. The solution was filtered through a Millipore syringe filter with 0.22 μm
203 porosity. The filtered extracts were then analyzed for both anions (SO₄²⁻, NO₃⁻, Cl⁻, NO₂⁻ and oxalate)
204 and cations (Na⁺, NH₄⁺, K⁺, Ca²⁺ and Mg²⁺) in parallel. The anions were analyzed using an anion self-
205 regenerating suppressor (ASRS 300) and an IonPac AS11-HC (2 x 250 mm) analytical column and
206 ~~measured~~ ~~detected~~ with a Dionex conductivity detector. For cations, a cation self-regenerating
207 suppressor (CSRS ULTRA II) and an IonPac CS18 (2 m x 250 mm) analytical column were used ~~in~~
208 ~~conjunction~~ ~~together~~ with a Dionex conductivity detector. The separation of anions was conducted using
209 25 mM KOH as an eluent at a flow rate of 0.38 ml/min, and the separation of cations was conducted
210 using 25 mM methanesulfonic acid at 0.25 ml/min.

211

212 The sum of nitrate and ammonium nitrogen ~~showed~~ ~~awas~~ ~~in~~ good agreement with ~~the~~ measured TN
213 (Fig. S1 in Supplementary Information (SI)), and based on the results of TN, NO₃⁻ and NH₄⁺, organic
214 nitrogen (OrgN) was also calculated using ~~the~~ following equation (Wang et al., 2010): OrgN = TN -
215 14*[NO₃⁻/62 + NH₄⁺/18].

216

2.5. EC/OC analysis

Online measurements of organic and elemental carbon (OC and EC) in aerosols were provided in parallel to the aerosol collection on quartz filters mentioned above by a field ~~semi-online~~ ~~semi-online~~ OC/EC analyzer (Sunset Laboratory Inc., USA) connected to a PM1 inlet. The instrument was equipped with a carbon parallel-plate denuder (Sunset Lab.) to remove volatile organic compounds to avoid a positive bias in the measured OC. Samples were taken at 4 h intervals, including the thermal-optical analysis, which lasts approximately 15 min. The analysis was performed using the shortened EUSAAR2 protocol: step [gas] temperature [°C]/duration [s]: He 200/90, He 300/90, He 450/90, He 650/135, He-Ox. 500/60, He-Ox. 550/60, He-Ox. 700/60, He-Ox. 850/100 (Cavalli et al., 2010). Automatic optical corrections for charring were made during each measurement, and a split point between EC and OC was detected automatically (software: RTCalc526, Sunset Lab.). Instrument blanks were measured once per day at midnight, and they represent only a background instrument ~~response~~ ~~signal~~ without ~~filter exposure~~ ~~any reflection on concentrations~~. Control calibrations using a sucrose solution were made before each change of the filter (ca. every 2nd week) to check the stability of instruments. The 24 h averages with identical measuring times, such as on quartz filters, were calculated from ~~the~~ acquired 4 h data. The sum of EC and OC provided ~~the~~ TC concentrations, which were consistent with ~~the~~ TC values measured by EA (see Fig. S2 in SI).

2.6. Spearman correlation calculations

Spearman correlation coefficients (r) were calculated using R statistical software (ver. 3.3.1). ~~The correlations~~ ~~Correlations~~ were calculated for the annual dataset (~~n=139~~ ~~and samples~~), separately for each season (autumn: ~~25~~, winter: ~~38~~, spring: ~~43~~, and summer: ~~33~~ ~~samples~~), and ~~the~~ winter ~~event~~ ~~Event~~ (7 ~~samples~~). Data from the winter *Event* were excluded from the annual and winter datasets for the correlation analysis as their distinctly high concentrations and isotopic values might have affected the results. Correlations with p-values over 0.05 were taken as statistically insignificant.

3. Results and discussion

The time series of TN, TC and their isotope ratios ($\delta^{15}\text{N}$ and $\delta^{13}\text{C}$) for the whole measurement campaign are depicted in Fig. 1. ~~Some sampling~~ ~~Sampling~~ gaps were caused in autumn and at the end of spring ~~are caused~~ by servicing or outages of the sampler. ~~However;~~ ~~however~~, 146 of the samples ~~from~~ ~~between~~ September 27, 2013, ~~to~~ ~~and~~ August 9, 2014, are ~~sufficient~~ ~~enough~~ for a seasonal study. In Fig. 1, the winter *Event* is highlighted, which has divergent values, especially for $\delta^{15}\text{N}$, and is discussed in detail in section 3.4.

253

254 Table 1 summarizes the results for ~~the~~ four seasons: autumn (Sep.—Nov.), winter (Dec.—Feb.),
255 spring (Mar.—May) and summer (Jun.—Aug.). The higher TN concentrations were observed in
256 spring (max. 7.59 $\mu\text{gN m}^{-3}$), while the higher TC concentrations were obtained during the winter *Event*
257 (max. 13.6 $\mu\text{gC m}^{-3}$). Conversely, the lowest TN and TC concentrations were observed in summer
258 (TabFig. 1).

259

260 Figure 2 shows ~~the~~ relationships between ~~the~~ TC and TN concentrations and their stable isotopes for
261 one year. The correlation between TC and TN is significant ($r=0.7170$), but ~~the relationship split~~ during
262 ~~high~~ concentration events, ~~this dependence can be split~~ due to ~~divergent source~~ the different
263 origins of these components. The highest correlations between TC and TN were obtained during
264 transition periods in autumn (0.85) and spring (0.80). Correlations between TC and TN in winter (0.43)
265 and summer (0.37) were weaker but still statistically significant ($p<0.05$). As seen in Table 1, the
266 seasonal averages of TC/TN ratios fluctuate, but their medians have similar values for autumn, winter
267 and spring. The summer TC/TN value is higher (3.45) and characteristic of a significant shift in
268 chemical composition, which is in line with previous studies at the site (Schwarz et al., 2016). However,
269 seasonal differences in the TC/TN ratios were ~~Correlations between TC and TN in winter (0.43) and in~~
270 ~~summer (0.37) were weaker but still statistically significant ($p<0.05$). As seen in the TC/TN ratios~~
271 ~~(Table 1), seasonal TC/TN averages fluctuate, but their medians have similar values for autumn, winter~~
272 ~~and spring, while the summer value is higher (3.45) and roughly points to different aerosol composition~~
273 ~~in comparison with other seasons. However, seasonal differences between TC/TN ratios are~~ not as large
274 as those in other works (e.g., Agnihotri et al., 2011), and thus, this ratio itself ~~did~~ does not provide much
275 information about aerosol sources.

276

277 The correlation between $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$ (Fig. 2, right) is also significant but negative (-0.6974). However,
278 there is a statistically significant correlation for spring only (-0.54), while in other seasons, correlations
279 are statistically insignificant. ~~(autumn: 0.29, winter: 0.11 and summer: 0.07)~~. This result highlights
280 ashows that significant ~~shift and related changes~~ in the sources of carbonaceous aerosols and their
281 isotope values ~~isotopic composition of nitrogen together with carbon occur especially in spring, while~~
282 ~~the sources were rather~~ there are stable sources of particles during other seasons, winter and summer.
283 The winter *Event* measurements show the highest $\delta^{13}\text{C}$ ~~values~~ and the lowest $\delta^{15}\text{N}$ values, but a they are
284 still in line with the linear fit does not show a significant differences as compared to rest of the fitting
285 ~~of all annual~~ data (Fig. 2, right).

286

287 3.1. Total nitrogen and its $\delta^{15}\text{N}$

288

289 The $\delta^{15}\text{N}$ values are stable in winter at approximately 15‰, with ~~an~~the exception of ~~the~~winter *Event*,
 290 which ~~showed~~~~deviated by~~ an average of 13‰. In summer, the $\delta^{15}\text{N}$ shows strong enrichment of ^{15}N in
 291 comparison with winter, resulting in an average value of 25‰. During the spring period, we
 292 ~~observed~~~~observe~~ a slow increase in $\delta^{15}\text{N}$ from April to June (Fig. 1), indicating a gradual change in
 293 nitrogen chemistry in the atmosphere. During autumn, a gradual change is not obvious because of a
 294 lack of data in a continuous time series. ~~The range of~~Year round, $\delta^{15}\text{N}$ ~~was~~ranged from 0.6‰ to 28.2‰
 295 ~~year round.~~‰. Such a ~~wid~~large range may ~~arise~~originate from ~~a~~the limited number of ~~nitrogen-main~~
 296 ~~compounds~~containing ~~species and/or components~~nitrogen in aerosols, which ~~are~~is specifically present
 297 in the ~~forms~~form of NO_3^- , NH_4^+ and/or organic nitrogen (OrgN), ~~and thus, the final $\delta^{15}\text{N}$ value in TN~~
 298 ~~can be formulated by the following equation:~~

$$299 \delta^{15}\text{N}_{\text{TN}} = \delta^{15}\text{N}_{\text{NO}_3^-} \cdot f_{\text{NO}_3^-} + \delta^{15}\text{N}_{\text{NH}_4^+} \cdot f_{\text{NH}_4^+} + \delta^{15}\text{N}_{\text{OrgN}} \cdot f_{\text{OrgN}}$$

300 ~~where $f_{\text{NO}_3^-} + f_{\text{NH}_4^+} + f_{\text{OrgN}} = 1$ and f represents the fractions of nitrogen from NO_3^- , NH_4^+ and OrgN in~~
 301 ~~TN, respectively).~~ The highest portion of nitrogen is contained in NH_4^+ (54 % of TN year-round),
 302 followed by OrgN (27 %) and NO_3^- (19 %). ~~Although~~While the NH_4^+ content in TN is seasonally stable
 303 (51-58 %, Table 1), the NO_3^- content is seasonally dependent; ~~the highest~~—higher in winter, ~~and~~
 304 ~~somewhat lower~~similarly balanced in spring and autumn. ~~In,~~and very low in summer; when the
 305 dissociation of NH_4NO_3 plays an important role ~~the NO_3^- content is very low,~~ and its nitrogen is
 306 partitioned from the aerosol phase to ~~the~~ gas phase (Stelson et al., 1979).

307
 308 The seasonal trend of $\delta^{15}\text{N}$ ~~of~~in TN, with the lowest values in winter and highest in summer, has been
 309 observed in other studies from urban Paris (Widory, 2007), rural Brazil (Martinelli et al., 2002), East
 310 Asian Jeju Island (Kundu et al., 2010) and rural Baengnyeong Island (Park et al., 2018) sites in Korea.
 311 However, different seasonal trends of $\delta^{15}\text{N}$ ~~of~~in TN in Seoul (Park et al., 2018) show that such seasonal
 312 variation does not always occur.

313
 314 Figure 3 shows changes in $\delta^{15}\text{N}$ values as a function of the main nitrogen components in TN, with
 315 different colors for different days. There are two visible trends for a type of nitrogen. Although ^{15}N is
 316 more depleted with increasing contents of NO_3^- in TN, the opposite is true for NH_4^+ and OrgN. The
 317 strongest dependence ~~for~~of most ~~of the~~ bulk data is expressed by a strong negative correlation between
 318 $\delta^{15}\text{N}$ and the ~~fraction~~share of NO_3^- in TN (Fig. 3). In all cases, the dependence during the winter *Event*
 319 is completely opposite to the rest of the bulk data (Fig. 3), ~~suggesting the presence of~~showing different
 320 processes ~~for~~on $\delta^{15}\text{N}$ ~~values~~formation, which is ~~characterized~~highlighted by a ~~very~~ strong positive
 321 correlation between $\delta^{15}\text{N}$ and NO_3^- -N/TN (0.98). This point will be discussed in section 3.4.

322
 323 Considering the individual nitrogen components, several studies (Freyer, 1991; Kundu et al., 2010;
 324 Yeatman et al., 2001b) ~~showed~~show seasonal trends of $\delta^{15}\text{N}$ of NO_3^- , with the lowest $\delta^{15}\text{N}$ in summer
 325 and the highest in winter. ~~Savard et al. (2017 and references therein)~~Savard et al. (2017 and references

326 ~~therein~~) summarized four possible reasons for this seasonality of $\delta^{15}\text{N}$ ~~of~~ NO_3^- : ~~namely, that is,~~ (i)
327 changes in NO_x emissions ~~strength~~, (ii) influence of wind directions in the relative contributions from
328 sources with different isotopic ~~composition~~ ~~composition~~, (iii) the effect of temperature on isotopic
329 fractionation and (iv) chemical transformations of nitrogen oxides over time with a lower intensity of
330 sunlight, which can lead to higher $\delta^{15}\text{N}$ values of atmospheric nitrate during winter months, as shown
331 by Walters et al. (2015a). In ~~the case of our study, it is most likely that data, mixing of all of these~~
332 factors ~~contributed, to a certain extent, to the nitrogen probably had an influence on the nitrate~~ isotopic
333 composition ~~of NO_3^- throughout during different parts of~~ the year.

334
335 Conversely, Kundu et al. (2010) reported higher $\delta^{15}\text{N}$ values of NH_4^+ in summer than in winter and
336 ~~generally~~ reported higher $\delta^{15}\text{N}$ values ~~of~~ NH_4^+ than ~~in~~ NO_3^- , except for winter ~~season~~. In sum, the
337 contribution of NH_4^+ to $\delta^{15}\text{N}$ overwhelms ~~that the contribution~~ of NO_3^- ~~to $\delta^{15}\text{N}$~~ . Additionally, TN is
338 composed of NH_4^+ , NO_3^- and OrgN. In ~~Figure Fig.~~ 3, we can observe ~~an~~ the enrichment of ^{15}N in TN in
339 summer when the lowest NO_3^- contribution occurs. Thus, higher $\delta^{15}\text{N}$ values of ~~$\delta^{15}\text{N}$ in~~ TN in summer
340 are mainly caused by ~~higher abundances of~~ NH_4^+ originating from $(\text{NH}_4)_2\text{SO}_4$, OrgN and ammonium
341 salts of organic acids.

342
343 Furthermore, ~~in summer,~~ we observed one of the largest enrichments of ^{15}N ~~of~~ TN ~~in summer~~ aerosols
344 ~~as compared to previous in comparison with other~~ studies (Kundu et al., 2010 and references therein),
345 which may be ~~explained by due to~~ several reasons. First, the ~~previous studies works mentioned above~~
346 mainly ~~focused on studied~~ total suspended particles (TSP); ~~aerosols~~; however, we ~~focused focus~~ on the
347 fine ~~PM1~~ fraction (PM_{10}), whose surface, ~~which~~ should be more reactive ~~due to a larger surface area per~~
348 ~~unit of aerosol mass~~ than the coarse fraction and consequently result in a higher abundance of ^{15}N during
349 the gas/particle partitioning ~~between~~ NH_3 and NH_4^+ . Second, ~~the fine accumulation aerosol fraction of~~
350 ~~the Aitken mode particles have persists for~~ a longer ~~residence period of~~ time in the atmosphere than the
351 coarse ~~mode~~ fraction, which is also a factor ~~that results in an leading to higher ^{15}N enrichment of ^{15}N~~ .
352 Indeed, Mkoma et al. (2014) reported average higher $\delta^{15}\text{N}$ ~~of~~ TN in fine ~~aerosols~~ (17.4‰, $\text{PM}_{2.5}$)
353 ~~than in comparison with~~ coarse aerosols (12.1‰, PM_{10}); ~~and~~ Freyer (1991) also reported higher $\delta^{15}\text{N}$
354 ~~of~~ NO_3^- (4.2‰ to 8‰) in fine aerosols ($< 3.5 \mu\text{m}$) in comparison with the coarse mode (-1.4‰ to
355 5.5‰). Third, a shorter sampling ~~interval of our time in this~~ work (24 h) leads to ~~more chance to~~
356 ~~collect the collection of samples with~~ episodic ~~samples such as values (see the winter Event, which could~~
357 ~~not) that would~~ be ~~clearly detected due to~~ averaged (overlapped) ~~aerosols~~ over a longer ~~sampling~~
358 ~~period time resolution~~ (e.g., weekly samples).

359
360 Similarly, as in this study, the highest $\delta^{15}\text{N}$ values in TN were observed in a few studies from the Indian
361 region (Aggarwal et al., 2013; Bikkina et al., 2016; Pavuluri et al., 2010) where biomass burning is ~~the~~
362 common ~~source~~, and ambient temperatures are high. Therefore, in addition to the above reasons,

363 temperature also plays a significant role in ^{15}N enrichment. This point will be discussed in more detail
364 in section 3.3.

365

366 Figure 4 shows the $\delta^{15}\text{N}$ ~~of~~ TN as a function of NO_3^- concentration. Samples. The $\delta^{15}\text{N}$ shows a peak
367 at approximately 14.4‰ with the highest NO_3^- increasing nitrate concentrations ($>6 \mu\text{g m}^{-3}$, $n=5$) show
368 an average $\delta^{15}\text{N}$ of $13.3 \pm 0.7\text{‰}$. Assuming that NO_3^- in the fine aerosol fraction consists predominantly
369 of NH_4NO_3 (Harrison and Pio, 1983), it can be stated that ammonium nitrate at the Košetice site is a
370 source of nitrogen at the Košetice site, with $\delta^{15}\text{N}$ values at approximately 13.34‰, which is similar to
371 the winter values of $\delta^{15}\text{N}$ in NO_3^- in other studies. Specifically, Kundu et al. (2010) reported a winter
372 average ~~value~~ of $\delta^{15}\text{N}$ ~~of~~ NO_3^- at +15.9 ‰ from a Pacific marine site at Gosan Island, South Korea,
373 whereas Freyer (1991) reported +9.2‰ in a moderately polluted site from Jülich, Germany. Yeatman
374 et al. (2001) reported approximately +9‰ from a Weybourne coastal site, UK. Park et al. (2018)
375 reported 11.9‰ in Seoul and 11.7‰ from a rural site in Baengnyeong Island, Korea.

376

377 Considering the $\delta^{15}\text{N}$ of nitrogen oxides, which are common precursors of particulate nitrate, we can
378 see that the $\delta^{15}\text{N}$ of nitrogen oxides generated by coal combustion (Felix et al., 2012; +6 to +13‰,
379 Heaton, 1990) or biomass burning (+14‰, Felix et al., 2012) are in thea same range with our $\delta^{15}\text{N}$
380 during the period of enhanced concentrations of NO_3^- . These $\delta^{15}\text{N}$ values of nitrogen oxides are also
381 significantly higher than those from vehicular exhaust (-13 to -2‰ Heaton, 1990; -19 to +9‰ Walters
382 et al., 2015b) or biogenic soil (-48 to -19‰, Li and Wang, 2008). Because of the only slight difference
383 between above reported $\delta^{15}\text{N}$ of nitrogen oxides and our $\delta^{15}\text{N}$ of TN during maximal NO_3^- events, the
384 isotope composition is probably influenced by the process of kinetic isotopic fractionation in fossil fuel
385 combustion samples during heating season as referred by Cieżka et al. (2016) as one of three possible
386 processes. Thus, $\delta^{15}\text{N}$ values around 13.3‰ (Fig. Thus, $\delta^{15}\text{N}$ values of approximately 14‰ (Fig. 4) are
387 probably characteristic of fresh emissions from heating (both coal and biomass burning) because these
388 values are obtained during the domestic heating season.

389

390 The exponential curves in Fig. 4 represent a boundary in which ~~the~~ $\delta^{15}\text{N}$ values are migrating as a result
391 of the enrichment or depletion of ^{15}N , which is associated with the removal or loading of NO_3^- in
392 aerosols. These curves represent two opposite chemical processes, with a match at approximately
393 13.34‰, which showed a strong logarithmic correlation ($r=0.96$ during winter *Event*, green line, and
394 -0.81 for the rest of points, black line, Fig. S3). These results indicate a significant and different
395 mechanism by which nitrogen isotopic fractionation occurs in aerosols. In both cases, the decrease in
396 nitrate leads to exponential changes in the enrichment or depletion of ^{15}N from a value of
397 13.3~~approximately 14‰~~. In the case of enrichment, in addition to a higher proportion of NH_4^+ than
398 NO_3^- , the dissociation process of NH_4NO_3 can cause an increase in ^{15}N ~~of~~ TN during a period of
399 higher ambient temperatures, as hypothesized by Pavuluri et al. (2010).

400

401 OrgN has not been widely studied as compared to particulate NO_3^- and NH_4^+ , although it represents a
402 significant fraction of TN (e.g., Jickells et al., 2013; Neff et al., 2002; Pavuluri et al., 2015). Figure 5
403 shows the relationship between $\delta^{15}\text{N}$ of TN and OrgN. Organic nitrogen consists organic compounds
404 containing nitrogen in water soluble and insoluble fractions. The majority of samples have a
405 concentration range of 0.1-0.5 $\mu\text{g m}^{-3}$ (gray highlight in Fig. 5), which can be considered as background
406 OrgN at the Košetice site. During the domestic heating season with the highest concentrations of NO_3^-
407 and NH_4^+ , we can observe a significant increase in OrgN with $\delta^{15}\text{N}$ again at approximately 13.314%,
408 which implies that the isotopic composition of OrgN is determined by the same sourceprocess during
409 maximal NO_3^- concentrations, that is, emissions from domestic heating. In the case of emissions from
410 combustion, OrgN originates mainly from biomass burning (Jickells et al., 2013 and references therein),
411 and thus, elevated concentrations of OrgN (as well as together with high NO_3^- and NH_4^+ conc.) may
412 refer to this source. On the other hand, looking at the trend of OrgN/TN in dependence on $\delta^{15}\text{N}$ (Fig. 3),
413 it is more similar to the trend of NH_4^+ -N/TN than NO_3^- -N/TN. Thus, it can be considered assumed that
414 the changes in the $\delta^{15}\text{N}$ of OrgN in samples highlighted as a gray area in Fig. 5 are probably driven
415 more by the same changes in NH_4^+ particles, and especially in summer with elevated OrgN in TN (Table
416 1).

417

418 3.2. Total carbon and its $\delta^{13}\text{C}$

419

420 ~~The $\delta^{13}\text{C}$ of TC ranged between -25.4‰ and -28.9‰ (Fig. 6), which is similar but broader than the~~
421 ~~range reported at a rural background site in Vavilhill (southern Sweden, range -26.7 to -25.6‰,~~
422 ~~Martinsson et al. (2017)), urban Wrocław (Poland, range -27.6 to -25.3‰.~~ The $\delta^{13}\text{C}$ of TC ranged from
423 -28.9 to -25.4‰ (Fig. 6) and the lowest $\delta^{13}\text{C}$ we observed in field blank samples (mean -29.2‰, n=7),
424 indicating that the lowest summer values in particulate matter were close to gas phase values. Our $\delta^{13}\text{C}$
425 values are within the range reported for particulate TC (-29‰ to -15‰) as summarized by Gensch et
426 al. (2014). The lowest values are associated with fine particles after combustion and transport (Ancelet
427 et al., 2011; Widory, 2006) while the highest values are associated with the coarse fraction and
428 carbonate contribution (Kawamura et al., 2004). This broad range can be explained by the influence of
429 marine aerosols (Ceburnis et al., 2016), different anthropogenic sources (e.g., Widory et al., 2004), as
430 well as different distributions of C3 and C4 plants (Martinelli et al., 2002) resulting in different $\delta^{13}\text{C}$
431 values in the northern and southern hemispheres (Górka et al. (2014)), and different sites (urban, coastal,
432 forest) in Lithuania (East Europe, Masalaite et al., 2015, 2017) but similar to those published by Fisseha
433 et al. (2009) in Zurich. However, our $\delta^{13}\text{C}$ values are smaller than those reported for coastal TSP
434 aerosols from Okinawa (East Asia, range -24.2 to -19.5‰, Kunwar et al. (2016)) or rural Tanzania
435 (Central East Africa, range -26.1 to -20.6‰ in PM2.5, Mkoma et al. (2014)). In fact, similar or different

436 ~~$\delta^{13}\text{C}$ values are widely reported in the northern and southern hemispheres (Cachier, 1989), which can~~
437 ~~be explained by different distributions of C3 and C4 plants (Martinelli et al., 2002), the influence of~~
438 ~~marine aerosols (Ceburnis et al., 2016), as well as different anthropogenic sources (e.g., Widory et al.,~~
439 ~~2004). The $\delta^{13}\text{C}$ values at the Košetice site fall within the range common to other European sites. The~~
440 ~~$\delta^{13}\text{C}$ values are significantly smaller than those of $\delta^{15}\text{N}$ due to a higher number of carbonaceous~~
441 ~~compounds in the aerosol mixture whose isotope ratio overlaps each other. However, it is possible to~~
442 ~~distinguish lower $\delta^{13}\text{C}$ values in summer (Table 1), which may indicate a contribution from higher~~
443 ~~terrestrial plant emissions. Similarly, Martinsson et al. (2017) reported lower $\delta^{13}\text{C}$ values in summer in~~
444 ~~comparison with other seasons, which they explain by high biogenic aerosol contributions from C3~~
445 ~~plants.~~

446
447 ~~A comparison of $\delta^{13}\text{C}$ with TC in Fig. 6 shows an enhanced enrichment of ^{13}C at higher TC~~
448 ~~concentrations. The lowest $\delta^{13}\text{C}$ values were observed in field blank samples (mean -29.2% , $n=7$),~~
449 ~~indicating that the lowest summer values in particulate matter were close to gas phase values. The $\delta^{13}\text{C}$~~
450 ~~values at the Košetice site fall within the range common to other European sites. For example, a rural~~
451 ~~background site in Vavihill (southern Sweden, range -26.7 to -25.6% , Martinsson et al. (2017)), urban~~
452 ~~Wroclaw (Poland, range -27.6 to -25.3% , Górká et al. (2014)), different sites (urban, coastal, forest) in~~
453 ~~Lithuania (East Europe, Masalaite et al., 2015, 2017), as well as urban Zurich (Switzerland, Fisseha et~~
454 ~~al. (2009)).~~

455 ~~The range of TC $\delta^{13}\text{C}$ values is significantly narrower than that of TN $\delta^{15}\text{N}$ due to a higher number of~~
456 ~~carbonaceous components in the aerosol mixture whose isotope ratio overlaps one another. However,~~
457 ~~it is possible to distinguish lower $\delta^{13}\text{C}$ values in summer (Table 1), which may indicate a contribution~~
458 ~~from higher terrestrial plant emissions. Similarly, Martinsson et al. (2017) reported lower $\delta^{13}\text{C}$ values~~
459 ~~in summer in comparison with other seasons, which they explain by high biogenic aerosol contributions~~
460 ~~from C3 plants.~~

461 ~~A comparison of $\delta^{13}\text{C}$ with TC in Fig. 6 shows an enhanced enrichment of ^{13}C at higher TC~~
462 ~~concentrations. A similar dependence of $\delta^{13}\text{C}$ on the TC concentration was observed by Fisseha et al.~~
463 ~~(2009), whereby winter ^{13}C enrichment was associated with WSOC (water soluble organic~~
464 ~~carbon) that originated mainly from wood combustion. Similarly, at the Košetice station, different~~
465 ~~carbonaceous aerosols were observed during the heating season (Oct.–Apr.) than in summer (Mbengue~~
466 ~~et al., 2018; Vodička et al., 2015). Moreover, whereby winter aerosols at the Košetice site were~~
467 ~~probably affected by not only biomass burning but also burning of coal (Schwarz~~
468 ~~et al., 2016), which can result in higher carbon contents and more ^{13}C -enriched particles (Widory,~~
469 ~~2006). However, relatively low $\delta^{13}\text{C}$ values in our range (up to -28.9%) are caused by not only sources~~
470 ~~of TC but also the fact that fine particles are more ^{13}C depleted in comparison with coarse~~

471 ~~particles~~ Furthermore, based on the number of size distribution measurements at the Košetice site, larger
472 ~~particles were observed in winter in comparison with summer, even in the fine particle fraction (Zíková~~
473 ~~and Ždímal, 2013), which can also have an effect on lower $\delta^{13}\text{C}$ values in summer. Thus, the relatively~~
474 ~~low $\delta^{13}\text{C}$ values in our range (up to -28.9‰) are because fine particles have lower $\delta^{13}\text{C}$ values in~~
475 ~~comparison with coarse particles probably due to different sources of TC. (e.g., Masalaite et al., 2015;~~
476 ~~Skipityté et al., 2016). Furthermore, based on the number of size distribution measurements at the~~
477 ~~Košetice site, larger particles were observed in winter in comparison with summer, even in the fine~~
478 ~~particle fraction (Zíková and Ždímal, 2013), which can also have an effect on lower $\delta^{13}\text{C}$ values in~~
479 ~~summer.~~

481 3.3. Temperature dependence and correlations of $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ with other variables

482
483 Tables 2 and 3 show Spearman's correlation coefficients (r) of $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ with different variables
484 that may reflect some effects on ~~isotope distributions~~ ~~these isotopes~~. In addition to year-round
485 correlations, correlations for each season, as well as for the *Event*, are presented separately.

486
487 Correlations of $\delta^{15}\text{N}$ in winter and summer are often opposite (~~see~~ e.g., for TN -0.40 in winter vs. 0.36
488 in summer, for NH_4^+ -0.42 in winter vs. 0.40 in summer), indicating that ~~changes in~~ aerosol chemistry
489 at the nitrogen level ~~is~~ ~~are~~ different in these seasons. Similarly, the contradictory dependence between
490 $\delta^{15}\text{N}$ and TN in summer and winter was observed by Widory (2007) ~~in PM10 samples from Paris.~~
491 ~~Widory (2007) connected this result with different primary nitrogen origin (road-traffic emissions in~~
492 ~~summer and no specific source in winter) and following secondary processes associated with isotope~~
493 ~~fractionation during degradation of atmospheric NOx leading to two distinct pathways for~~ ~~on PM10~~
494 ~~samples from Paris and was connected with secondary processes affecting the nitrogen chemistry that~~
495 ~~follows two distinct pathways between~~ ^{15}N enrichment (summer) and depletion (winter).

496
497 From a meteorological point of view, a significant correlation of $\delta^{15}\text{N}$ with temperature has been
498 obtained, indicating the influence of temperature on the nitrogen isotopic composition. ~~The~~
499 ~~dependence~~ ~~Dependence~~ of $\delta^{15}\text{N}$ ~~of~~ ~~in~~ TN on temperature (Fig. 7) is ~~similar to that observed by Cieżka~~
500 ~~et al. (2016) for $\delta^{15}\text{N}$ of NH_4^+ from precipitation; however, it is the~~ opposite ~~of~~ ~~to~~ that observed by Freyer
501 (1991) for $\delta^{15}\text{N}$ ~~of~~ ~~in~~ NO_3^- . ~~The aforementioned studies; however, it is same to that observed by Cieżka~~
502 ~~et al. (2016) for $\delta^{15}\text{N}$ in NH_4^+ from precipitation. These authors~~ concluded that the isotope equilibrium
503 exchange between nitrogen oxides and particulate nitrates is temperature dependent and could lead to
504 more ^{15}N enriched NO_3^- during the cold season (Freyer et al., 1993; Savard et al., 2017). Although
505 Savard et al. (2017) reported a similar negative ~~temperature dependence for~~ $\delta^{15}\text{N}$ ~~of~~ ~~in~~ NH_4^+ ~~dependence~~
506 ~~at temperatures~~ in Alberta (Canada), ~~such as for~~ NO_3^- , most studies ~~reported a positive temperature~~

507 dependence for $\delta^{15}\text{N}$ of NH_4^+ that is stronger than that for $\delta^{15}\text{N}$ of NO_3^- (e.g., Kawashima and Kurahashi,
508 2011; Kundu et al., 2010). The reason is that reported the opposite temperature dependence for $\delta^{15}\text{N}$ in
509 NH_4^+ because the NH_3 gas concentrations are highermore abundant during warmerwarm weather
510 conditions, and thus the isotopic equilibrium exchange reaction, i.e., $\text{NH}_3(\text{g}) \leftrightarrow \text{NH}_4^+(\text{p})$, which leads
511 leading to ^{15}N enrichment in particles, is more intensive.

512
513 All the considerations mentioned above indicate that a resultingfinal relationship between $\delta^{15}\text{N}$ ofin TN
514 and temperature is driven by the prevailing nitrogen species, which is NH_4^+ in our case. A similar
515 dependence was reported by Pavuluri et al. (2010) between temperature and $\delta^{15}\text{N}$ ofin TN in Chennai
516 (India), where NH_4^+ strongly prevailed. They found the best correlation between $\delta^{15}\text{N}$ and temperature
517 during the colder period (range 18.4-24.5°C, avg. 21.2°C); however, during warmer periods, this
518 dependence was weakened. In our study, we observed the highest correlation of $\delta^{15}\text{N}$ with temperature
519 in autumn ($r=0.58$, temp. range -1.9 to 13.9°C, avg. 6.6°C), followed by spring ($r=0.52$, temp. range
520 1.5-18.7°C, avg. 9.3°C), but there was even a negative but insignificant correlation in summer ($r=-0.21$,
521 temp. range: 11.8-25.5°C, avg. 17.7°C). This result indicates that ambient temperature plays an
522 important role in the enrichment/depletion of ^{15}N ; however, it is not determined by a specific
523 temperature range but rather the conditions for repeating the process of “evaporation/condensation”,
524 as shown by the comparison with the work of Pavuluri et al. (2010). It is likely that isotopic fractionation
525 caused by the equilibrium reaction of $\text{NH}_3(\text{g}) \leftrightarrow \text{NH}_4^+(\text{p})$ reaches a certain level of enrichment under
526 higher temperature conditions in summer.

527
528 In summer, $\delta^{15}\text{N}$ correlates positively with NH_4^+ ($r=0.40$) and SO_4^{2-} (0.51), indicating a link with
529 $(\text{NH}_4)_2\text{SO}_4$ that is enriched by ^{15}N due to aging. Figure 8 shows an enrichment of ^{15}N as a function of
530 the molar ratio of $\text{NH}_4^+/\text{SO}_4^{2-}$. The highest $\text{NH}_4^+/\text{SO}_4^{2-}$ ratios, showing an ammonia rich atmosphere,
531 were observed during winter, late autumn and early spring along with high abundance of NO_3^- that is
532 related to favorable thermodynamic conditions during heating season and enough ammonia in the
533 atmosphere. Gradual decreasing molar ratios of $\text{NH}_4^+/\text{SO}_4^{2-}$ during spring indicate a gradual increase of
534 ambient temperatures and therefore worsened thermodynamic conditions for NO_3^- formation in aerosol
535 phase, which was accompanied by a visible decrease in the nitrate content in aerosols (Fig. 8). The
536 increase of temperatures finally leads to the $\text{NH}_4^+/\text{SO}_4^{2-}$ ratio reaching 2 at the turn of spring and summer.
537 Finally, summer values of $\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratio below 2 indicate that SO_4^{2-} in aerosol particles at high
538 summer temperatures may not be completely saturated with ammonium but it can be composed from
539 mixture of $(\text{NH}_4)_2\text{SO}_4$ and NH_4HSO_4 (Weber et al., 2016). The equilibrium reaction between these two
540 forms of ammonium sulfates related to temperature oscillation during a day and due to vertical mixing
541 of the atmosphere is a probable factor which leads to increased values of $\delta^{15}\text{N}$ in early summer.
542 Ammonia measurements, that were carried out at the Košetice site until 2001, showed that NH_3
543 concentrations in summer were slightly higher than in winter

544 (http://portal.chmi.cz/files/portal/docs/uoco/isko/tab_roc/2000_enh/CZE/kap_18/kap_18_026.html),
545 which supports temperature as a main factor influencing $\text{NH}_4^+/\text{SO}_4^{2-}$ ratio at Košetice. In this context,
546 we noticed that 25 out of 33 summer samples have molar ratios of $\text{NH}_4^+/\text{SO}_4^{2-}$ below 2, and the
547 remaining samples are approximately 2, and the relative abundance of NO_3^- in PM1 in those samples is
548 very low (ca. 1.7 %). ~~Figure 8 shows a decreasing molar ratio of $\text{NH}_4^+/\text{SO}_4^{2-}$ with increasing $\delta^{15}\text{N}$~~
549 ~~enrichment, especially during spring, indicating a gradual uptake of ammonia in the gas phase to aerosol~~
550 ~~phase. With a decreasing $\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratio, there is also a visible decrease in the nitrate content~~
551 ~~in aerosols (Fig. 8). However, when the $\text{NH}_4^+/\text{SO}_4^{2-}$ ratio approaches a value below 2, there is not~~
552 ~~enough available ammonia in the gas phase, leading to the exclusion of nitrate from the aerosol phase,~~
553 ~~as well as to the disruption of the thermodynamic equilibrium between $\text{NH}_3(\text{g}) \leftrightarrow \text{NH}_4^+(\text{p})$, which~~
554 ~~previously led to $\delta^{15}\text{N}$ enrichment in the particles. In this context, we note that 25 out of 33 summer~~
555 ~~samples have molar $\text{NH}_4^+/\text{SO}_4^{2-}$ ratios below 2, and the remaining samples are approximately 2,~~
556 ~~although the average relative abundance of NO_3^- in PM1 in those samples is very low (ca. 1.7 %).~~

557
558 Recently, Silvern et al. (2017) reported that organic aerosols can play a role in modifying or retarding
559 the achievement of $\text{H}_2\text{SO}_4\text{-NH}_3$ thermodynamic equilibrium at $\text{NH}_4^+/\text{SO}_4^{2-}$ molar ratios of less than 2,
560 even when sufficient amounts of ammonia are present in ~~the~~ gas phase. Thus, an interaction between
561 sulfates and ammonia may be hindered due to the preferential reactions such that organics coated with
562 aged aerosols coated with organics preferentially react (Liggio et al., 2011). In thermodynamic
563 equilibrium, partitioning between gas (NH_3) and aerosol (NH_4^+) phases should result in even larger $\delta^{15}\text{N}$
564 values of particles in summer, however, measurements show a different situation. Summer $\delta^{15}\text{N}$ values
565 are highest but further enrichment was stopped. Moreover ~~Indeed~~, we observed a positive (and
566 significant) correlation between temperature and $\delta^{13}\text{C}$ ($r=0.39$) only in summer, whereas the correlation
567 coefficient of $\delta^{15}\text{N}$ vs. temperature is statistically insignificant, negative (-0.21), suggesting that while
568 values of $\delta^{15}\text{N}$ reached their maxima, the $\delta^{13}\text{C}$ can still grow with even higher temperatures due to the
569 ~~thermodynamic equilibrium between $\text{NH}_3(\text{g})$ and nitrogen in particles was minimal or replaced by the~~
570 influence of organics in summer ~~this~~ season. ~~Ammonia measurements directly at the Košetice site were~~
571 ~~carried out until 2001, and they showed that the NH_3 concentrations in summer and winter were~~
572 ~~comparable~~
573 (http://portal.chmi.cz/files/portal/docs/uoco/isko/tab_roc/2000_enh/CZE/kap_18/kap_18_026.html),
574 which ~~indirectly support the above hypothesis.~~

575
576 As seen in Table 3, summertime ~~The summer~~ positive correlations of $\delta^{13}\text{C}$ with ozone ($r=0.66$) and
577 temperature (0.39) indicate oxidation processes that can indirectly lead to an enrichment of ^{13}C in
578 organic aerosols that are enriched with oxalic acid (Pavuluri and Kawamura, 2016). ~~carbon isotope~~
579 ~~enrichment~~. This result is also supported by the fact that the content of oxalate in PM1, measured by
580 IC, was twice as high in spring and summer than in winter and autumn. The influence of temperature

581 on $\delta^{13}\text{C}$ in winter is opposite to that in ~~the~~ summer. The ~~winter~~-negative correlation (-0.35) in winter
582 probably ~~indicates points to the evolution of~~ more fresh emissions from domestic heating (probably coal
583 burning) with higher $\delta^{13}\text{C}$ values ~~contents of ^{13}C~~ during cold season ~~lower temperatures~~.

584
585 The whole year temperature dependence on $\delta^{13}\text{C}$ is the opposite of that observed for $\delta^{15}\text{N}$ (Fig. 7, left),
586 suggesting more ^{13}C -depleted products in summer. This result is probably connected with different
587 carbonaceous aerosols during winter (anthropogenic emissions from coal, wood and biomass burning
588 with the enrichment of ^{13}C) in comparison with the summer season (primary biogenic and secondary
589 organic aerosols with lower $\delta^{13}\text{C}$) (Vodička et al., 2015). ~~This result is probably connected with different~~
590 ~~carbonaceous aerosols during winter (anthropogenic emissions from coal, wood and biomass burning~~
591 ~~with the enrichment of ^{13}C) in comparison with the summer season (primary biogenic and secondary~~
592 ~~organic aerosols with lower $\delta^{13}\text{C}$).~~ The data of $\delta^{13}\text{C}$ in Fig. 7 are also more scattered, which indicates
593 that in the case of carbon, the isotopic composition depends more on sources than on temperature.

594
595 Correlations of $\delta^{13}\text{C}$ with OC are significant in all seasons; they are strongest in spring and weakest in
596 summer (Table 3). Correlations of $\delta^{13}\text{C}$ with EC, whose main sources are ~~source is~~ combustion processes
597 from domestic heating and transportation, are significant ($r=0.61-0.88$) only during the heating season
598 (autumn–spring, see Table 3), while in summer, the correlation is statistically insignificant (0.28). Thus,
599 the isotopic composition of aerosol carbon at the Košetice station is not significantly influenced by EC
600 emitted from transportation; otherwise the year-round correlation between $\delta^{13}\text{C}$ and EC would suggest
601 that transportation is also ~~be~~ significant source of EC in summer. This result can be biased by the fact
602 that EC constitutes on average 19% of TC during all seasons. However, it ~~This result~~ is consistent with
603 positive correlations between $\delta^{13}\text{C}$ and gaseous NO_2 , as well as particulate nitrate, which is also
604 significant ~~in~~ from autumn to spring. This, and this result is also supported by the negative correlation
605 of $\delta^{13}\text{C}$ with the EC/TC ratio ($r=-0.51$), which is significant only in summer.

606
607 It should be mentioned that the wind directions during the campaign were similar, with the exception
608 of ~~the~~ winter season, when southeast (SE) winds prevailed (see Fig. S4 in SI). We did not observe any
609 specific dependence of isotopic values on wind directions, except for the *Event*.

610 3.4. Winter Event

611
612 The winter *Event* represents ~~the~~ a period ~~from~~ between January 23 ~~to~~ and February 5, 2014, when an
613 enrichment of ^{13}C and substantial depletion of ^{15}N occurred in PM1 (see Figs. 1 and 9 for details). We
614 do not observe any trends of the isotopic compositions of $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ with wind directions, except
615 for the period of the *Event* and one single measurement on ~~18th~~ December 18, 2013. Both the *Event* and
616 the single measurement are connected to SE winds through Vienna and the Balkan Peninsula (Fig. 10).

617 More elevated wind speeds with very stable SE winds are observed on the site with samples showing
618 the most ^{15}N depleted values at the end of the *Event* (Fig. 9). Stable weather conditions and the
619 homogeneity of the results indicate a local or regional source, which is probably associated with the
620 formationemissions of sulfates (Fig. S5), ~~which are not sufficiently mixed at this time.~~

621
622 Although the *Event* contains only 7 samples, high correlations are obtained for $\delta^{15}\text{N}$ and $\delta^{13}\text{C}$ (Tables 2
623 and 3). Generally, correlations of $\delta^{15}\text{N}$ with several parameters during the *Event* are opposite to those
624 of ~~the~~ four seasons, indicating the exceptional nature of these aerosols from a chemical point of view.
625 During the *Event*, $\delta^{15}\text{N}$ correlates positively with NO_3^- ($r=0.96$) and NO_3^- -N/TN (0.98). Before the *Event*,
626 we also observed the highest), with large values of $\delta^{15}\text{N}$ at approximately 13.314‰, which we
627 previously interpreted as an influence of the emissions from domestic heating viaby coal and/or biomass
628 burning. Positive correlations of $\delta^{13}\text{C}$ with oxalate and potassium (both 0.93) and the negative
629 correlation with temperature (-0.79) also suggestshow that the *Event* is associated with fresh emissions
630 from burning sourcecombustion.

631
632 In contrast, we find that most $\delta^{15}\text{N}$ values with a depletion of ^{15}N are associated with enhanced NH_4^+
633 contents (70-80 % of TN) and ~~the~~ almost-total absence of NO_3^- nitrogen (see Figs. 3 and 4). Although
634 some content of OrgN is detected during the *Event* (Fig. 3), the correlation between $\delta^{15}\text{N}$ and OrgN/TN
635 is not significant (Table 2). This result suggestsshows that nitrogen with the lowest $\delta^{15}\text{N}$ values is
636 mainly connected with NH_4^+ , which is supported by a strong negative correlation between $\delta^{15}\text{N}$ and
637 NH_4^+ /TN (-0.86). Assuming that nitrogen in particles mainly originates from gaseous nitrogen
638 precursors via gas-to-particle conversion (e.g., Wang et al., 2017) during the *Event*, we couldshould
639 expect the nitrogen originatedto originate mainly from NH_3 with depleted ^{15}N but not nitrogen oxides.
640 Agricultural emissions from both fertilizer application and animal waste are importantsueh sources of
641 NH_3 -emissions (Felix et al., 2013). Considering possible agriculture emission sources, there exist
642 several collective farms, with both livestock (mainly cows, HolsteinHolsteins cattle) and crop
643 production in the SE direction from the Košetice observatory – namely, Agropodnik Košetice (~~in~~ 3.4
644 km awaydistance), Agrodám Hořepník (6.8 km) and Agrošev Červená Řečice (9.5 km). Skipitytė et al.
645 (2016) reported lower $\delta^{15}\text{N}$ values of TN (+1 to +6‰) for agriculture-derived particulate matter of
646 poultry farms, which are close to our values obtained during the *Event* (Fig. 9). -

647
648 The $\delta^{15}\text{N}$ values from the *Event* are associated with an average temperature of below 0°C (Figs. 7 and
649 9). Savard et al. (2017) observed the lowest values of $\delta^{15}\text{N}$ ofin NH_3 with temperatures below -5°C , and
650 the NH_4^+ particles that were simultaneously sampled were also isotopically lighter compared to the
651 samples collected under higher temperature conditions. They interpreted the result as a preferential dry
652 deposition of heavier isotopic $^{15}\text{NH}_3$ species during the cold period, whereas lighter $^{14}\text{NH}_3$ species
653 preferentially remains in the atmosphere. However, cold weather can also lead to a decline of ammonia

654 fluxes from aerosol water surfaces, soil, etc. (Roelle and Aneja, 2002), which generally result in a deficit
655 of ammonia in the atmosphere. Emissions from farms are not as limited by low temperature and are
656 thus a main source of ammonia in this deficiency state. The removal of NH₃ leads to a non-equilibrium
657 state between the gas and aerosol phases. They interpreted this result as the preferential dry deposition
658 of heavier isotopic ¹⁵NH₃ species during the cold period, whereas the remaining lighter ¹⁴NH₃ species
659 in the atmosphere, lead to lighter NH₄⁺ in particles. Moreover, the removal of NH₃ by dry deposition
660 also leads to a non-equilibrium state between the gas and aerosol phases. Such an absence of equilibrium
661 exchange of NH₃ between the gas and liquid/solid phases is considered to cause the supported by a
662 NH₄⁺/SO₄²⁻ molar ratiosratio below 2 for the three most ¹⁵N depleted samples (Fig. 8). However, underIn
663 such conditions, nitrate partitioning in PM is negligible. It should be mentioned, that a deficiency of
664 ammonia in atmosphere during the winter Event leads to completely opposite δ¹⁵N values than in
665 summer (see section 3.3) even if molar ratios NH₄⁺/SO₄²⁻ are below 2 in both cases.

666
667 Unidirectional, and unidirectional reactions of isotopically lighter NH₃ isotope with H₂SO₄ in the
668 atmosphere are strongly preferred bydue to the kinetic isotope effect, which is -(after several minutes,)
669 followed by enrichment of ¹⁴NH₃the nitrogen due to the newly established equilibrium (Heaton et al.,
670 1997). Based on laboratory experiments, Heaton et al. (1997) estimated the isotopic enrichment factor
671 between gas NH₃ and particle NH₄⁺, ε_{NH₄-NH₃}, to be +33‰. Savard et al. (2017) reported an isotopic
672 difference (Δδ¹⁵N) between NH₃ (g) and particulate NH₄⁺ as a function of temperature, whereas Δδ¹⁵N
673 for a temperature of approximately 0°C was approximately 40‰. In both cases, after subtraction of
674 these values (33 or 40‰) from the δ¹⁵N values of the measured *Event*, we obtain values frombetween
675 approximately -40 to -28‰, which are in a range of δ¹⁵N-NH₃ (g) measured for agricultural emissions.
676 These values are especially in good agreement with δ¹⁵N of NH₃ derived from cow waste (ca. -38 to -
677 22‰, Felix et al., 2013).

678
679 Thus, during the coursein case of the winter Event, we probably observedobserve PM representing a
680 mixture of aerosols from household heating characterized by higher amounts of NO₃⁻ and low value
681 (8.2‰) of δ¹⁵N ofin TN₂ (ca. 14‰), which are gradually replaced by ¹⁵N-depleted agricultural aerosols.
682 TheResults of the whole process occurred under low temperature conditionsfrom low temperatures that
683 was first initiated by a deficiency support dry deposition of NH₃ followed by an unidirectional (kinetic)
684 reaction of isotopically lighter isotope-NH₃(g) → NH₄⁺(p), in which NH₃ is) originate mainly originated
685 from agricultural sources in the SE ofdirection from the Košetice station.

686
687 If the four lowest values of δ¹⁵N mainly represent agricultural aerosols, then it can be suggested that the
688 δ¹³C values from the same samples should originate from samealso be characteristic of agricultural
689 sources. During the winter Event, In this case, the δ¹³C values ranging from -26.2 to -25.4‰ belong to
690 the most ¹³C enriched fine aerosols at the Košetice site. However, similar δ¹³C values were reported by

691 Widory (2006) for particles from coal combustion (-25.6 to -24.6‰). Skipitytė et al. (2016) reported a
692 mean value of $\delta^{13}\text{C}$ ~~ofin~~ TC (-23.7±1.3‰) for PM1 particles collected on a poultry farm, and ~~they~~
693 suggested the litter as a possible source for the particles. Thus, in the case of $\delta^{13}\text{C}$ values that we
694 observed during the winter Event are probably caused by we observed, emissions ~~either~~ from domestic
695 heating ~~than from and/or~~ agricultural sources. This is also supported by increased emissions of SO₂ from
696 coal combustion to formation of sulfates, are responsible for the ¹³C values.
697

698 4. Summary and Conclusions

699
700 Based on the analysis of year-round data of stable carbon and nitrogen isotopes, we extracted ~~some~~
701 important information on the processes taking place in fine aerosols during different seasons at the
702 Central European station of Košetice. Seasonal variations were observed for $\delta^{13}\text{C}$ and $\delta^{15}\text{N}$, as well as
703 for TC and TN concentrations. The supporting data (i.e., ions, EC/OC, meteorology, trace gases)
704 revealed characteristic processes that led to changes in the isotopic compositions on the site.

705 The main and gradual changes in nitrogen isotopic composition occurred in spring. During early spring,
706 domestic heating with wood stoves is still common, with high nitrate concentrations in aerosols, which
707 decreased toward the end of spring. Additionally, the temperature slowly increases, and the overall
708 situation leads to thermodynamic equilibrium exchange between gas (NO_xNO_{x3}-NH₃-SO₂ mixture) and
709 aerosol (NO₃⁻- NH₄⁺- SO₄²⁻ mixture) phases, which causes an¹⁵N enrichment of ¹⁵N in aerosols.
710 Enrichment of ¹⁵N ($\Delta\delta^{15}\text{N}$) from the beginning to the end of spring was approximately +10‰. Gradual
711 springtime changes in isotopic composition were also observed for $\delta^{13}\text{C}$, but the depletion was small,
712 and $\Delta\delta^{13}\text{C}$ was only -1.4‰.

713
714 In summer, we observed the lowest concentrations of TC and TN; however, there was an enhanced
715 enrichment of ¹⁵N, which was probably caused by the aging of ~~nitrogenous nitrogen~~ aerosols, where
716 ammonium sulfate and bisulfate is subjected to isotopic fractionation via equilibrium exchange between
717 NH₃(g) and NH₄⁺(p) when. ~~Based on a NH₄⁺/SO₄²⁻ molar ratio was of less than 2, ~~we concluded that~~
718 ~~summer aerosols become more acidic, and thus, kinetic isotopic fractionation took place via the~~
719 ~~equilibrium exchange of nitrogen species~~. However, summer values of $\delta^{15}\text{N}$ were still among the highest
720 compared with those in previous studies, which can be explained by several factors. First, a fine aerosol
721 fraction (PM1) is more reactive, and its residence time in the atmosphere is longer than coarse mode
722 particles, leading to ¹⁵N enrichment in aged aerosols. Second, summer aerosols, compared to other
723 seasons, contain a negligible amount of nitrate, contributing to a decrease in the average value of $\delta^{15}\text{N}$
724 of TN. Although the summer $\delta^{15}\text{N}$ values were the highest further ¹⁵N enrichment was minimized at
725 this season. On the other hand, we observed an enrichment of ¹³C only in summer, which can be
726 explained by the photooxidation processes of organics and is supported by the positive correlation of~~

727 $\delta^{13}\text{C}$ with temperature and ozone. Despite this slow enrichment process, summertime $\delta^{13}\text{C}$ values were
728 the lowest compared to those in other seasons and referred predominantly to organic aerosols of
729 biogenic origin. ~~The role of organics in summer may also have an effect on the aforementioned ^{15}N~~
730 ~~enrichment due to thermodynamic equilibrium.~~

731
732 In winter, we found the highest concentrations of TC and TN. Lower winter $\delta^{15}\text{N}$ values were apparently
733 influenced by fresh aerosols from combustion, which were strongly driven by the amount of nitrates
734 (mainly NH_4NO_3 in PM1), and led to an average winter value ($13.3 \pm 0.7\%$) of $\delta^{15}\text{N}$ of
735 ~~TN, approximately 14%.~~ Winter $\delta^{13}\text{C}$ values were more enriched than summer values, which are
736 involved with the and they were connected mainly to emissions from coal and (mostly) biomass and
737 coal burning for domestic heating.

738
739 We observed an aerosol event in winter, which was characterized by low temperatures below the
740 freezing point, stable southeast winds, and a unique isotope signature with a depletion of ^{15}N and
741 enrichment of ^{13}C . The winter *Event* characterized by ^{15}N depletion was probably caused by
742 ~~preferential the dry deposition of NH_3 (with heavier isotope) during cold weather, and with decreasing~~
743 ~~concentrations of NO_3^- . However, it was completely opposite to a summertime decrease in nitrate,~~
744 ~~which led to an enrichment of ^{15}N . In the case of the most depleted ^{15}N event, nitrate was suppressed to~~
745 ~~partition in aerosol and gas phases with~~ unidirectional reactions between of isotopically light ammonia,
746 and sulfuric acid resulting in $(\text{NH}_4)_2\text{SO}_4$, which originated mainly from agriculture emissions, and
747 sulfuric acid, resulting in $(\text{NH}_4)_2\text{SO}_4$ and NH_4HSO_4 . This process was probably supported by long-term
748 cold weather leading to a deficiency of ammonia in the atmosphere (due to dry deposition and/or low
749 fluxes), and subsequent suppression of nitrate to partitioning in aerosol in this case.

750
751 The majority of the yearly data showed a strong correlation between $\delta^{15}\text{N}$ and ambient temperature,
752 demonstrating an enrichment of ^{15}N via isotopic equilibrium exchange between the gas and particulate
753 phases. This process seemed to be one of the main mechanisms for ^{15}N enrichment at the Košetice site,
754 especially during spring. The most ^{15}N -enriched summer and most ^{15}N -depleted winter samples were
755 limited for by the partitioning of nitrate ~~in aerosols and suppressed equilibrium exchange~~ between
756 gaseous NH_3 and aerosols, aerosol NH_4^+ .

757
758 This study revealed a picture of the seasonal cycle of $\delta^{15}\text{N}$ in aerosol TN at the Košetice site. ~~The~~ In the
759 ~~case of carbon, the~~ seasonal cycle of $\delta^{13}\text{C}$ cycle values was not so pronounced because they mainly
760 depend on the isotopic composition of primary sources, which often overlapped. Although
761 photochemical, and because secondary oxidation reactions are driven were influenced by the kinetic
762 isotopic effect, the while phase transfer probably did not play a crucial role in the case of carbon at the
763 Central European site.

764

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766

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1015 doi:10.4209/aaqr.2013.02.0056, 2013.

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1019 Table 1: Seasonal and entire campaign averages \pm standard deviations, (medians in brackets) of
 1020 different variables.

	Autumn	Winter	Spring	Summer	Year
N of samples	25	45	43	33	146
TC [$\mu\text{g m}^{-3}$] (from EA)	3.61 \pm 1.61 (3.30)	4.76 \pm 2.44 (3.88)	3.78 \pm 2.03 (3.04)	2.71 \pm 0.76 (2.68)	3.81 \pm 2.03 (3.35)
TN [$\mu\text{g m}^{-3}$]	1.56 \pm 1.18 (1.33)	1.67 \pm 0.96 (1.45)	2.00 \pm 1.62 (1.47)	0.81 \pm 0.29 (0.82)	1.56 \pm 1.22 (1.26)
$\delta^{13}\text{C}$ [‰]	-26.8 \pm 0.5 (-26.9)	-26.7 \pm 0.5 (-26.7)	-27.1 \pm 0.5 (-27.0)	-27.8 \pm 0.4 (-27.7)	-27.1 \pm 0.6 (-27.0)
$\delta^{15}\text{N}$ [‰]	17.1 \pm 2.4 (16.9)	13.1 \pm 4.5 (15.2)	17.6 \pm 3.5 (17.3)	25.0 \pm 1.6 (25.1)	17.8 \pm 5.5 (16.9)
TC/PM1 [%]	28 \pm 6 (26)	33 \pm 8 (32)	38 \pm 15 (35)	31 \pm 6 (30)	33 \pm 11 (31)
TN/PM1 [%]	11 \pm 3 (11)	11 \pm 3 (12)	17 \pm 4 (17)	9 \pm 2 (9)	12 \pm 4 (12)
NO₃⁻-N/TN [%]	21 \pm 6 (21)	25 \pm 8 (28)	22 \pm 8 (21)	5 \pm 3 (4)	19 \pm 10 (20)
NH₄⁺-N/TN [%]	51 \pm 6 (51)	51 \pm 9 (49)	58 \pm 7 (60)	57 \pm 6 (57)	54 \pm 8 (54)
OrgN/TN [%]	28 \pm 8 (26)	25 \pm 8 (23)	20 \pm 8 (19)	39 \pm 6 (38)	27 \pm 10 (25)
TC/TN	2.77 \pm 1.10 (2.60)	3.34 \pm 1.66 (2.68)	2.33 \pm 0.98 (2.34)	3.60 \pm 1.23 (3.45)	3.01 \pm 1.38 (2.61)

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1022 Table 2: Spearman correlation coefficients (r) of $\delta^{15}\text{N}$ with various tracers. Only bold values are
 1023 statistically significant (p-values < 0.05).

$\delta^{15}\text{N}$ vs.	Autumn	Winter*	Spring	Summer	Year*	Event
TN	-0.30	-0.40	-0.70	0.36	-0.54	0.93
TN/PM1	-0.63	-0.50	-0.02	0.37	-0.35	0.36
NO₃⁻-N/TN	-0.39	-0.04	-0.73	-0.26	-0.77	0.98
NH₄⁺-N/TN	0.16	-0.30	0.60	0.52	0.42	-0.86
OrgN/TN	0.20	0.38	0.20	-0.33	0.51	-0.71
NO₃⁻	-0.41	-0.35	-0.80	-0.03	-0.78	0.96
NH₄⁺	-0.22	-0.42	-0.61	0.40	-0.44	0.75
OrgN	-0.26	-0.27	-0.56	0.30	-0.25	0.71
SO₄²⁻	-0.07	-0.38	-0.30	0.51	0.03	-0.57
Cl	-0.37	-0.18	-0.74	-0.37	-0.74	0.99
O₃ (gas)	0.45	0.14	0.15	-0.02	0.40	-0.71
NO₂ (gas)	-0.53	-0.34	-0.72	0.20	-0.64	0.86
NO₂/NO (gas)	-0.51	-0.26	-0.82	0.14	-0.76	0.82
Temp.	0.58	0.30	0.52	-0.21	0.77	-0.43

1024 *Event data are excluded from winter and year datasets.

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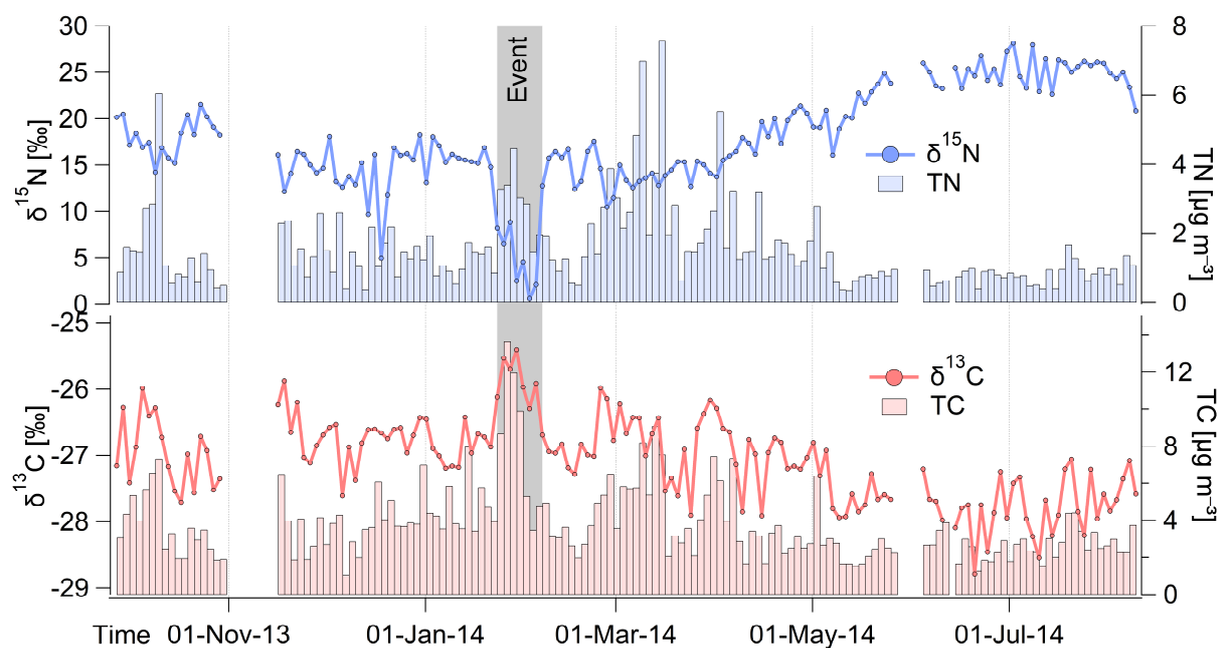
1030 Table 3: Spearman correlation coefficients (r) of $\delta^{13}\text{C}$ with various tracers. Only bold values are
 1031 statistically significant (p-values < 0.05).

$\delta^{13}\text{C}$ vs.	Autumn	Winter*	Spring	Summer	Year*	Event
OC	0.64	0.63	0.91	0.39	0.75	0.75
EC	0.61	0.74	0.88	0.28	0.84	0.46
EC/TC	0.06	0.06	0.13	-0.51	0.32	-0.32
TC/PM1	-0.16	-0.05	-0.40	0.22	-0.09	0.32
NO_3^-	0.74	0.52	0.71	0.12	0.76	0.39
NH_4^+	0.84	0.59	0.80	0.42	0.66	0.75
Oxalate	0.34	0.62	0.71	0.65	0.25	0.93
SO_4^{2-}	0.80	0.64	0.73	0.41	0.34	0.54
K^+	0.84	0.63	0.70	0.47	0.76	0.93
Cl^-	0.44	0.62	0.68	0.44	0.76	0.25
CO (gas)	0.21	0.53	0.60	0.32	0.37	0.68
O_3 (gas)	-0.41	-0.26	0.14	0.66	-0.33	0.11
NO_2 (gas)	0.67	0.38	0.70	0.18	0.69	0.32
NO_2/NO (gas)	0.72	0.65	0.67	0.68	0.78	0.96
Temp.	-0.33	-0.35	-0.20	0.39	-0.57	-0.79

1032 *Event data are excluded from winter and year datasets.

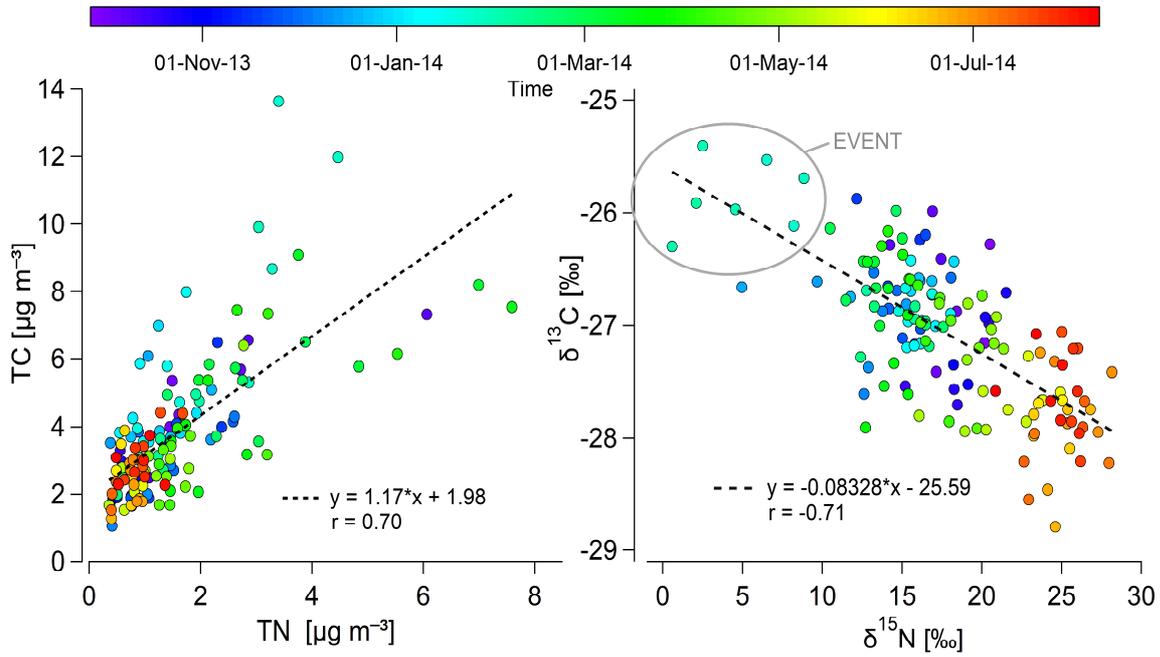
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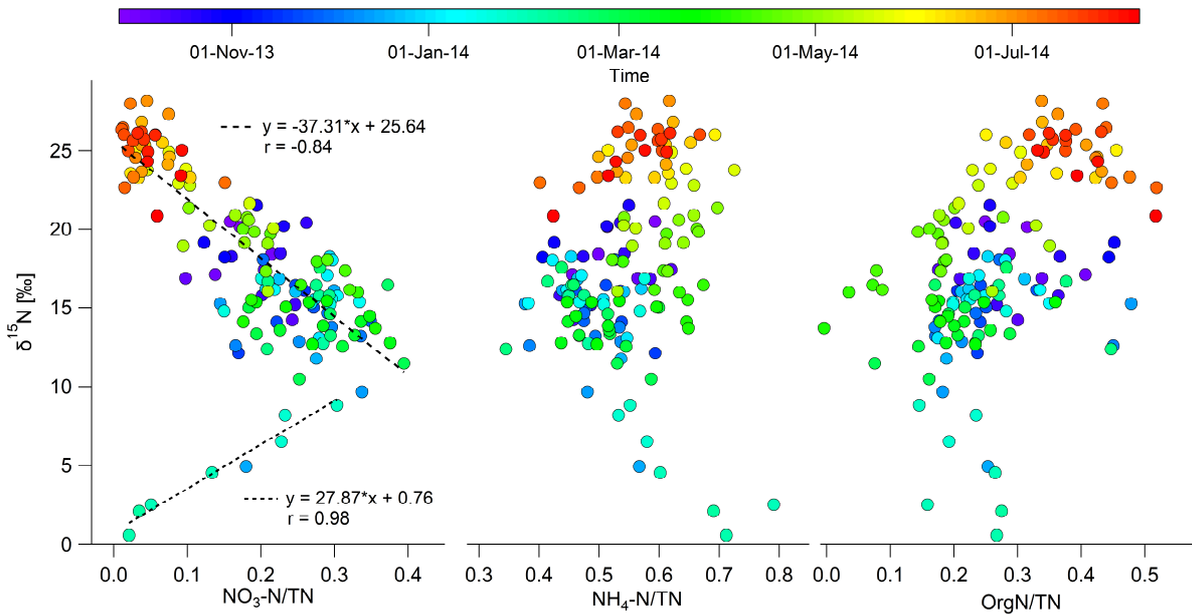
1036 Fig. 1: Time series of $\delta^{15}\text{N}$ ~~along together~~ with TN (top) and $\delta^{13}\text{C}$ ~~as well as together with~~ TC (bottom)
 1037 in PM1 aerosols at the Košice station. The gray color highlights an *Event* with divergent values,
 1038 especially for $\delta^{15}\text{N}$.



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1040 Fig. 2: Relationships between TC and TN (left) and their stable carbon and nitrogen isotopes (right).
 1041 The color scale reflects the time of sample collection. The gray circle highlights the winter *Event*
 1042 measurements.

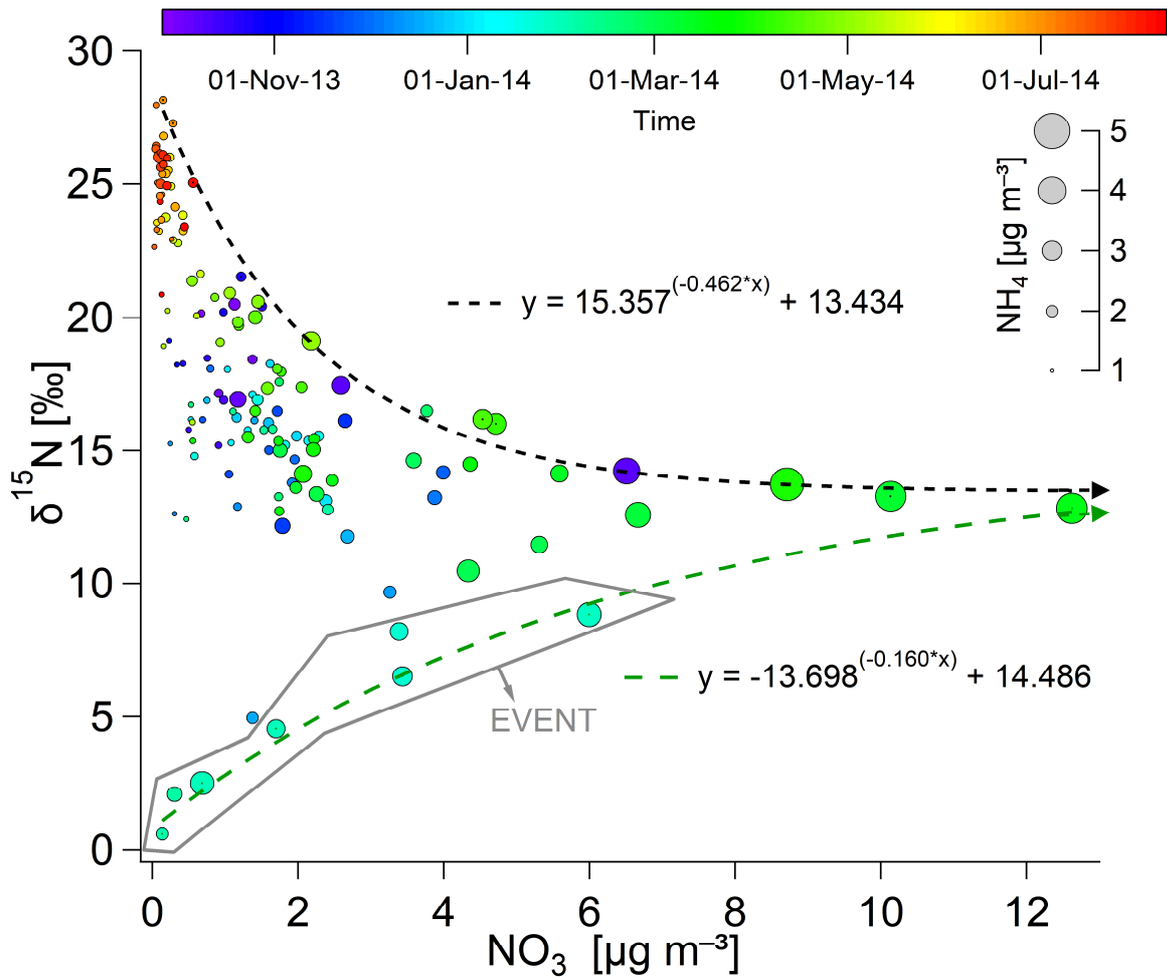
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1045 Fig. 3: Changes in $\delta^{15}\text{N}$ depending on fraction of individual nitrogen components ($\text{NO}_3\text{-N}$, $\text{NH}_4\text{-N}$, and
 1046 OrgN) in TN. The color scale reflects the time of sample collection.

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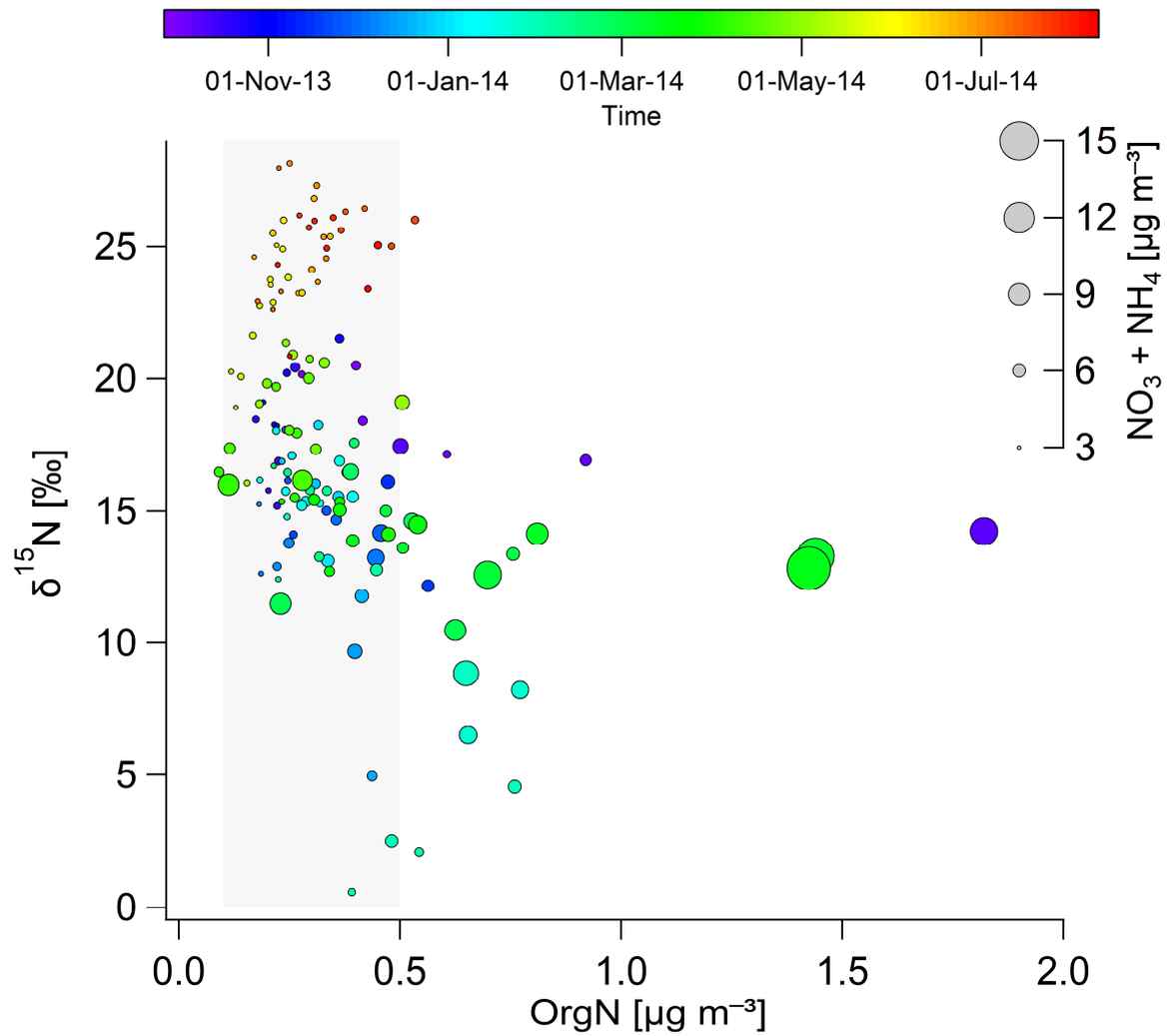


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1049 Fig. 4: Relationships of $\delta^{15}\text{N}$ of TN vs. NO_3^- concentrations. The larger circles indicate higher NH_4^+
 1050 concentrations. The color scale reflects the time of sample collection.

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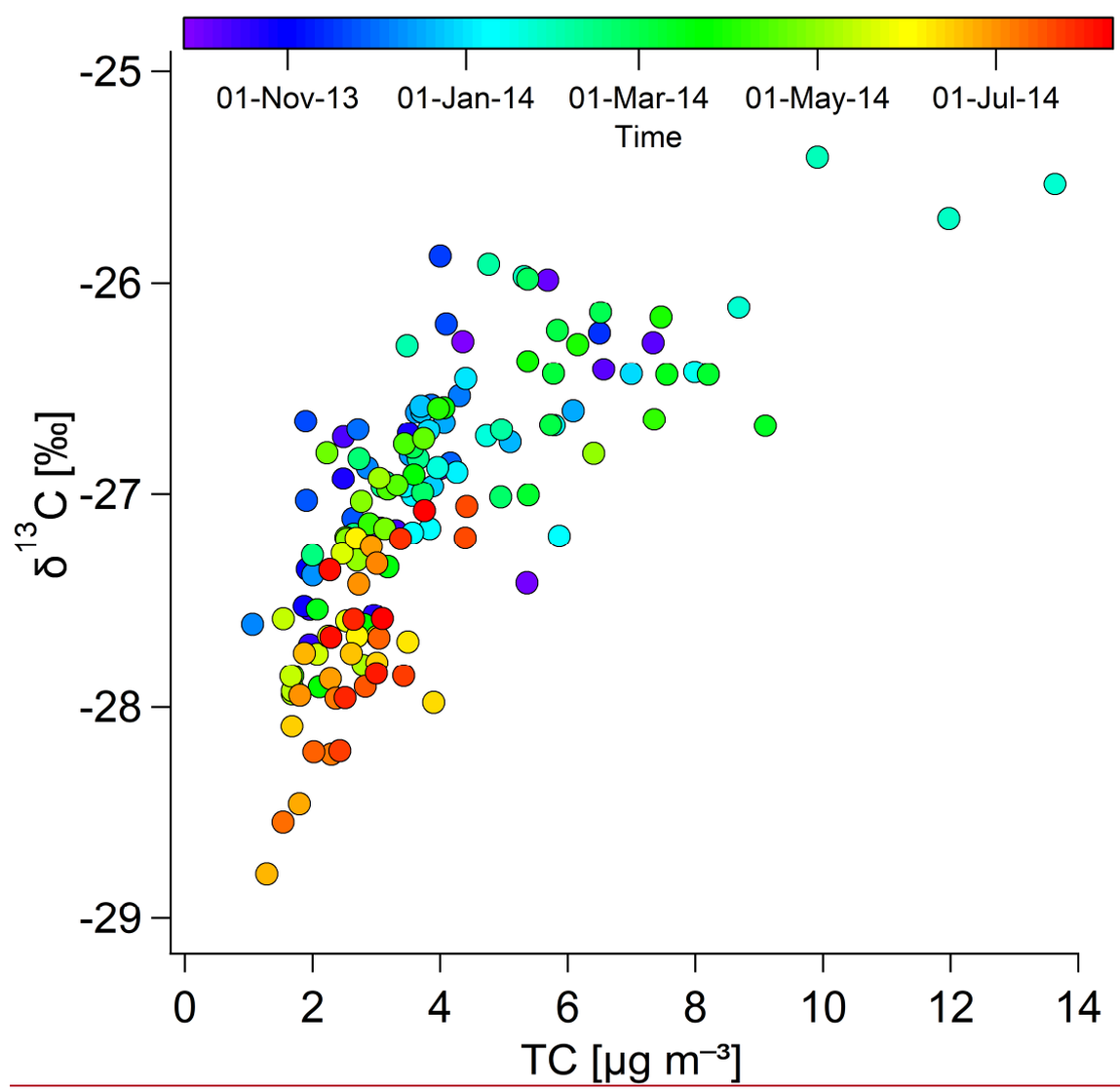
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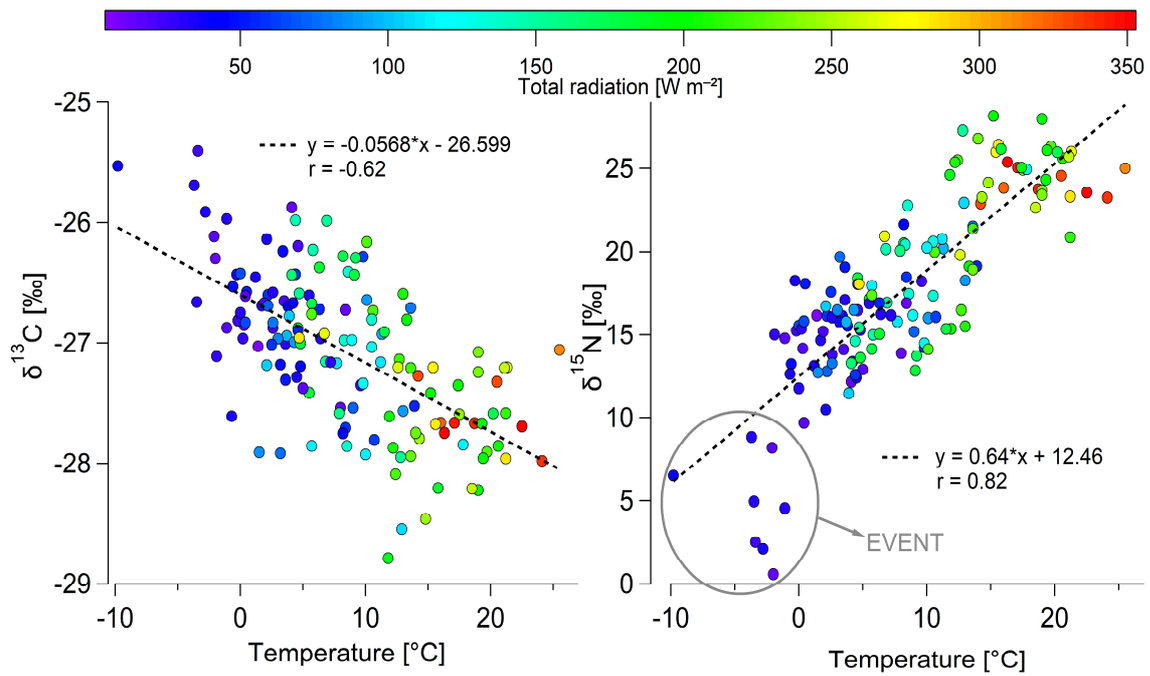
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Fig. 5: Relationships of $\delta^{15}\text{N}$ ~~of~~ TN vs. OrgN concentrations. The larger circles indicate higher sums of $\text{NO}_3^- + \text{NH}_4^+$ concentrations. The color scale reflects the time of sample collection, and the highlighted portion is a concentration range between $0.1\text{-}0.5 \mu\text{g m}^{-3}$.



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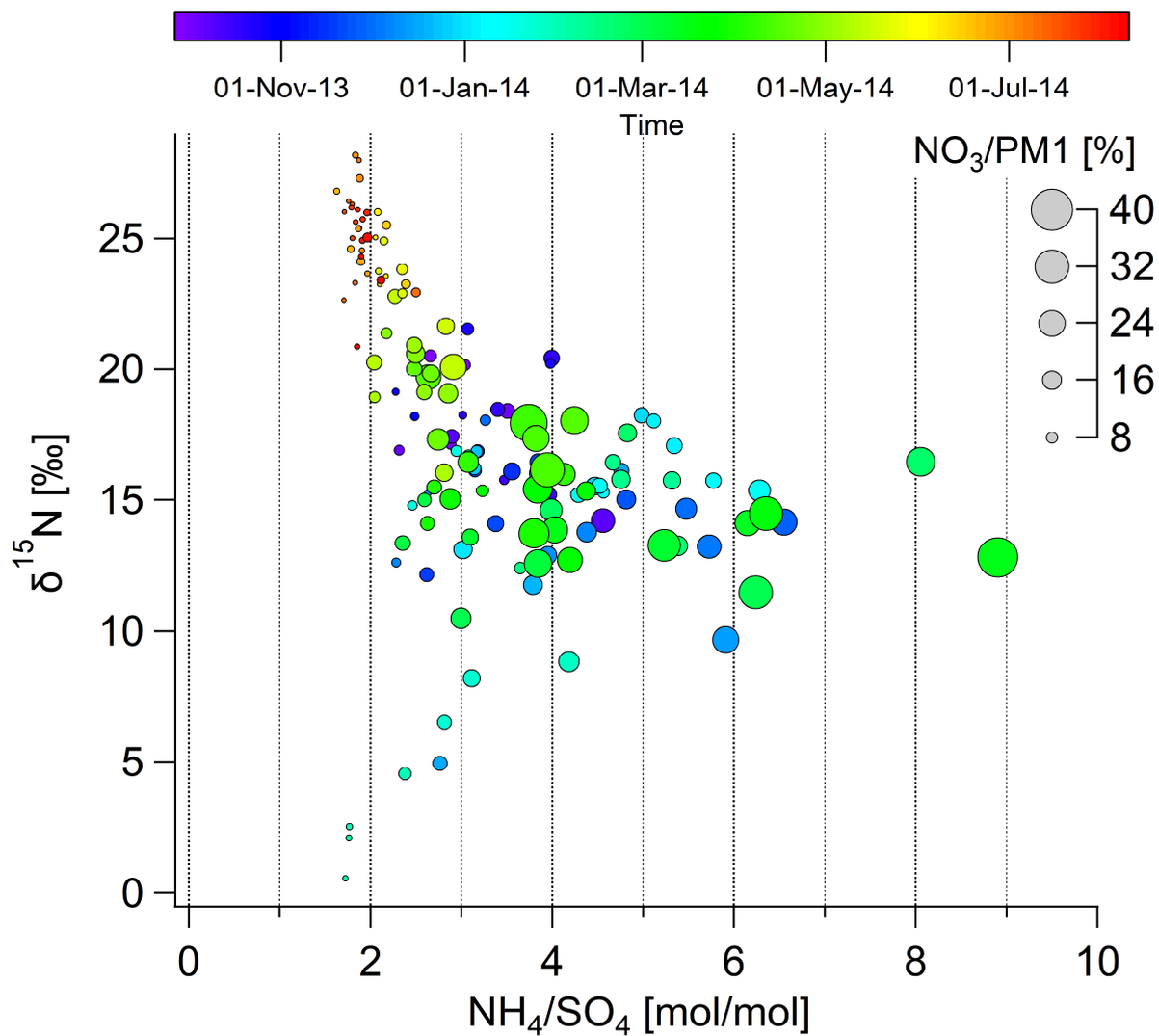
Fig. 6: Relationship between TC and $\delta^{13}\text{C}$. The color scale reflects the time of sample collection.



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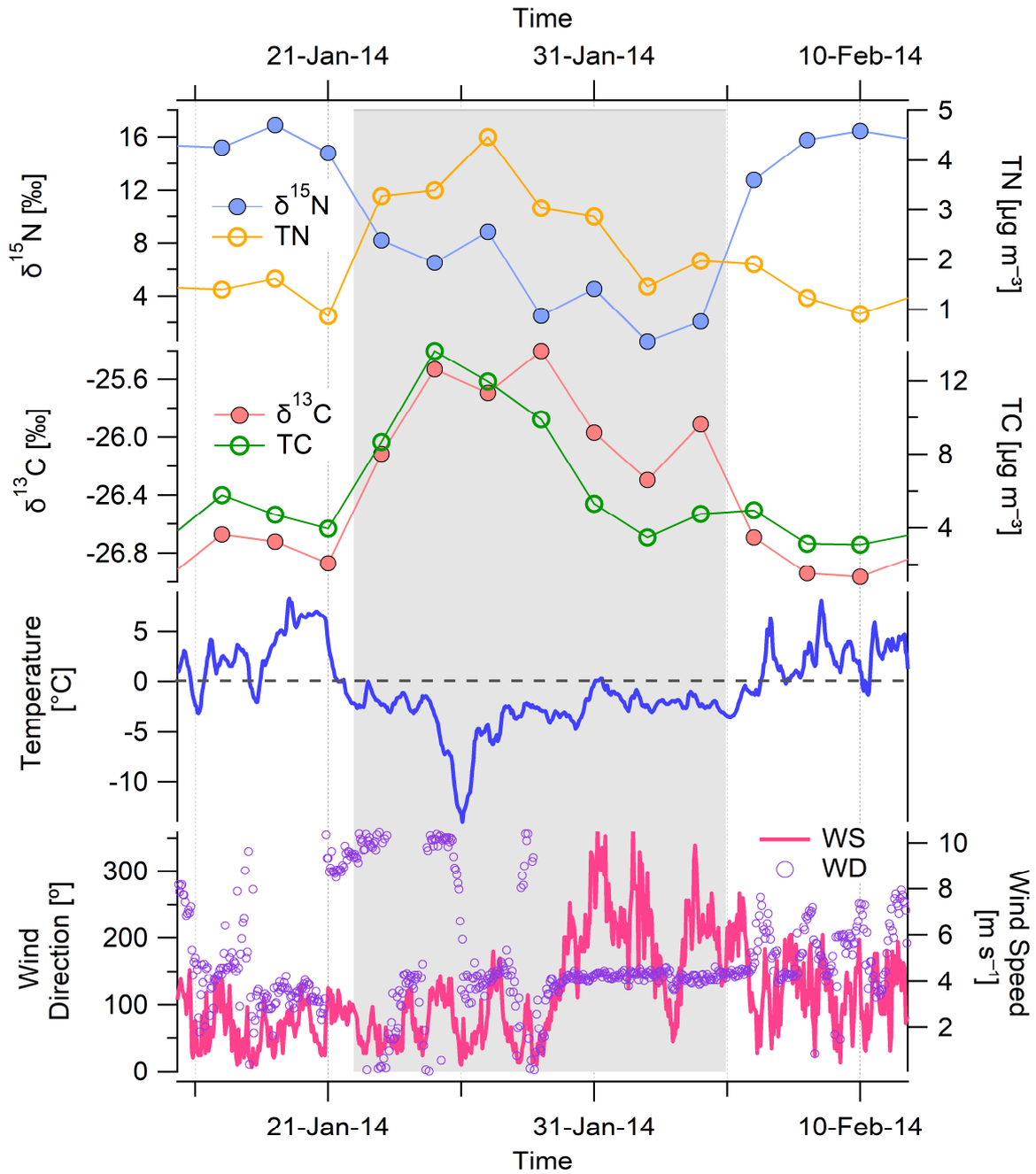
Fig. 7: Relationships between temperature and δ¹³C of TC (left) and δ¹⁵N of TN (right). The color scale reflects the total radiation.



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1064 Fig. 8: Relationships between $\delta^{15}\text{N}$ of TN and molar ratios of $\text{NH}_4^+/\text{SO}_4^{2-}$ in particles. The larger circle
 1065 indicates a higher nitrate content in PM1. The color scale reflects the time of sample collection.

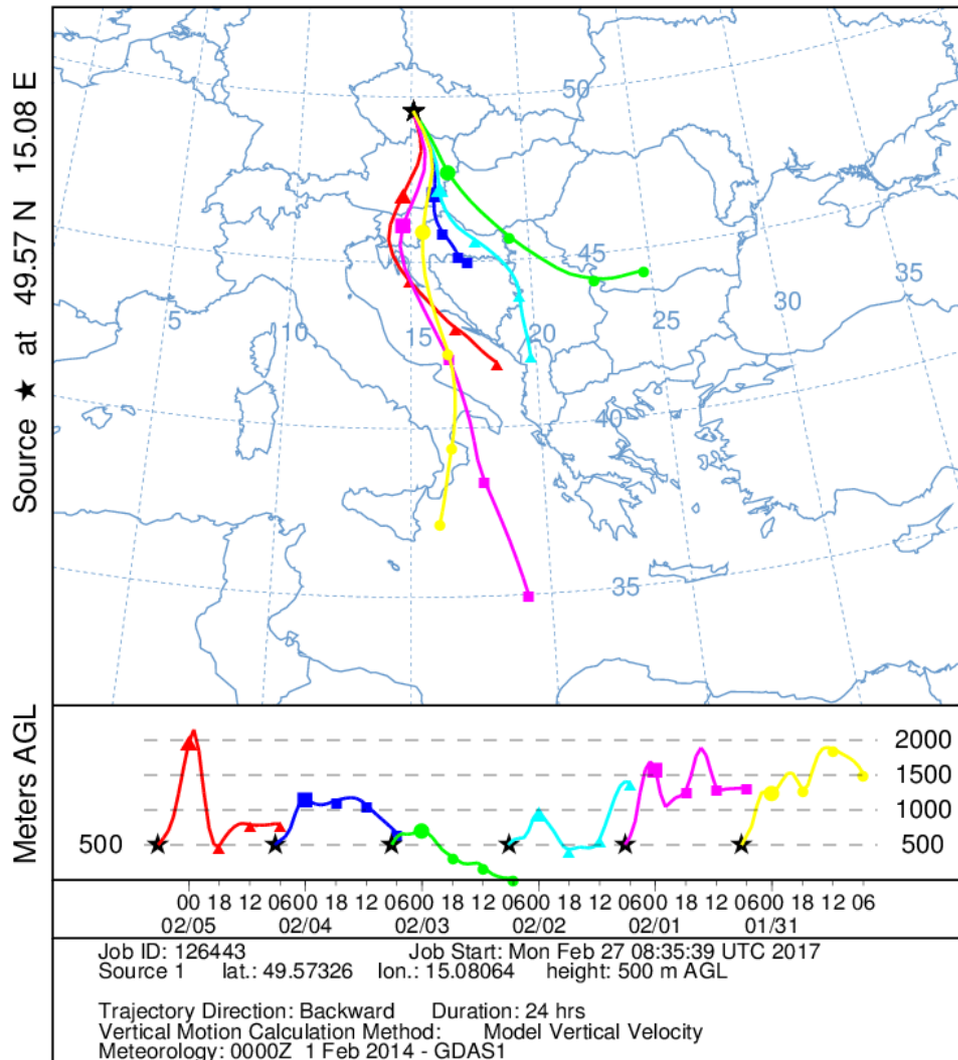
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Fig. 9: Time series of $\delta^{15}\text{N}$, TN, $\delta^{13}\text{C}$, TC and meteorological variables (temperature, wind speed and direction, 1 h time resolution) during the *Event*, which is highlighted by thea gray color.

NOAA HYSPLIT MODEL
 Backward trajectories ending at 0600 UTC 05 Feb 14
 GDAS Meteorological Data



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 1072 Fig. 10: NOAA HYSPLIT (Stein et al., 2015) 24 h backward air mass trajectories at 500 m above
 1073 ground level for the observation site from 30 Jan until 5 Feb 2014 (right).