

Referee's comments in black

Authors' responses in blue

## Responses to Tariq Munir (Referee 1)

### General comments

This manuscript is very well articulated, and it will attract readers working on plant PFT/traits responses to permafrost thaw and subsequent nutrient concentration in gelisols in the event of climate warming. The paper adds to our knowledge about how shrubs will respond to climate warming with their strategies to fight back by trading-off between traits for their sustained growth.

We thank Tariq Munir for taking the time to read and comment on our manuscript.

The manuscript does not describe how many times and how frequent the experimental sites were visited during the study years to imagine the field-work extent of this manuscript which tries to provide many solid conclusions. Without this information, it looks like the sites were set up and flowed by a couple of campaigns each year.

The experimental site was visited periodically during the study years as described in the reference we included in Line 101 (Wang *et al.*, 2017). The campaigns were limited to the growing season; due to the harsh conditions of the study area, the experimental plots were not accessible during the rest of the year. Below, we enumerate the visits to the plots (information extracted from Wang *et al.*, 2017).

The experimental plots were selected in July 2010; the heating cables were inserted in the soil also in this period. The following growing season (July 2011), the experimental treatments (heating and fertilization) were implemented in July 2011: the heating cables were connected to the solar panels in the heated plots and slow-release NPK fertilizer tablets were added to the fertilized plots. In 2013, nutrients were added again to the plots. During the experimental period, environmental factors were measured periodically. The permafrost thaw depth and soil moisture were measured 2-4 times per growing season, while soil temperature was measured continuously (data were recovered from the temperature loggers each growing season). Resin bags to assess soil nutrient availability were inserted in each plot in 2010 and replaced by new ones at the beginning of each August until 2014. The species abundance within the plots was recorded in 2010 and 2013 (results published in Wang *et al.*, 2017). Aboveground and belowground biomass was harvested in August 2014 (Wang *et al.*, 2017). The individuals selected for the plant trait analysis and growth rate (Iturrate-Garcia *et al.*, 2017) were sampled in the second half of 2014 growing season (31 July - 12 August 2014), when leaf and stem traits were also measured (indicated in Lines 128-129).

The above information will be added to section 2 Materials and Methods (subsections 2.2 Experimental design and 2.4 Study species and sampling).

Iturrate-Garcia, M., Heijmans, M.M.P.D., Schweingruber, F.H., Maximov, T.C., Niklaus, P.A., Schaepman-Strub, G. (2017). Shrub growth rate and bark responses to soil warming and nutrient addition – A dendroecological approach in a field experiment. *Dendrochronologia*, 45, 12-22.

Wang, P., Limpens, J., Mommer, I., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above and belowground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 164–175.

No pre-experimental conditions of the selected blocks are provided (if they were similar or different with some statistical analyses) which could be another drawback of this research.

Although no pre-experimental condition of the selected blocks are provided, potential differences within blocks were statistically considered by including "block" (factor with 5 levels) as fixed term in all the linear mixed-effect models. Moreover, in order to take into account species-specific trait differences among blocks, we added the

interaction between species and block (fix terms) to the statistical models, which we run for plant traits (detailed in subsection "2.6 Statistical analysis").

The statistical analyses/models performed might need a quick look back or rerun with random effects. There are no repeated measurements over the years I know of.

Except for the statistical models used to test the effects of treatments on soil temperature and permafrost thaw depth (see explanation at specific comment further down, referring to Line 166-170), the other statistical models (effects of treatments on plant traits) consider plot as random effect, as described in subsection "2.6 Statistical analysis". It is correct that there are no repeated measurements over the years.

### Specific comments

Paras 2-3: resource acquisition and conservative strategies of plants. do authors have references to these strategies studied

We will add the reference of Diaz *et al.* (2016) on Line 44, which refers to acquisitive and conservative strategies of plants.

Díaz, S., Kattge, J., Cornelissen, J.H.C., Wright, I.J., Lavorel, S., Dray, S. et al. (2016). The global spectrum of plant form and function, *Nature*, 529, 1-17.

Line 52. Would you like to put a reference for projected conditions?

We will add the article by Post *et al.* (2019) on Line 52, as a reference for projected conditions in the Arctic.

Post, E., Alley, R.B., Christensen, T.R., Macias-Fauria, M., Forbes, B.C., Gooseff, M.N., Iler, A., Kerby, J.T., Laidre, K.L., Mann, M.E., Olofsson, J., Stroeve, J.C., Ulmer, F., Virginia, R.A., Wang, M. (2019). The polar regions in a 2°C warmer world. *Science Advances*, 5, eaaw9883.

Line 54. I would better put a semi-colon here instead of two parentheses

OK

Line 62. Please correct referencing here I know, one can derive specific objectives from the last paragraph of hypothesis and an overview of the experimental components; however, I would better explicitly mention specific objectives helpful for researchers skimming several studies at a time

If we understood correctly, in this comment Taquir Munir (Referee 1) raises his concern about using solely the reference Violle *et al.* (2007) instead of adding several studies at a time. The inclusion of this reference, however, aims to clarify the concept of trait used in the studies related to tundra shrub responses to climate change. Violle *et al.* (2007) is a theoretical/review paper. This paper introduces the concept of "performance traits" and specifically elaborates on traits included in this category. Addressing the suggestion of the received Short Comment by Michael O'Brien (L62), we rephrased the sentence ('[...] performance traits (detailed in Violle *et al.*, 2007) [...]'), which might help to address also the comment by Taquir Munir.

In case that this comment refers to Line 74 instead of Line 62, we will rephrase the sentence to specifically mention the objectives of the study. 'The objective of this study is to experimentally investigate...'

Line97. Define growing season (e.g., May-Oct); I think, it must be a point when the daily maximum temperature reaches a minimum of 6-degree C. The growing season was never defined except table 1

We use the term "growing season" for the period of the year during which arctic plants photosynthesize and grow. The timing of the growing season depends on the air temperature, snowmelt, and thawing of frozen soil at the

study site. Air temperature (e.g. 6°C) alone is therefore not an appropriate proxy for the growing season and it can vary from year to year ( $\pm 4$  days; Parmentier *et al.* 2011) and from species to species. For the research area at the ecosystem level, the growing season lasts approximately from end of June to end of August, based on carbon flux analyses (Parmentier *et al.* 2011, van der Molen *et al.* 2007).

To address the referee's comment we will add an indication of time on Line94: '...and lichens. The growing season lasts from the end of June to end of August. The slightly acidic soil...'

Parmentier, F.J.W., van der Molen, M.K., van Huissteden, J., Karsanaev, S.A., Kononov, A.V., Suzdalov, D.A., Maximov, T.C., Dolman, A.J. (2011). Longer growing seasons do not increase net carbon uptake in the northeastern Siberian tundra. *Journal of Geophysical Research*, 116, G04013.

van der Molen, M.K.; van Huissteden, J., Parmentier, F.J.W., Petrescu, A.M.R., Dolman, A.J. *et al.* (2007). The growing season greenhouse gas balance of a continental tundra site in the Indigirka lowlands, NE Siberia. *Biogeosciences*, European Geosciences Union, 4 (6), pp. 985–1003.

Line102. What was the extent of a block? A schematic may help here

The six plots within each block were spaced by 1-2 meter distance from each other. With a size of 1.5 m x 1.5 m, all plots within a block were located in an area of approximately 10 m x 10 m. A picture of a typical block setup is contained in the supplementary material of Wang *et al.*, 2017. We will add the approximate extent of blocks to the experimental design section.

Wang, P., Limpens, J., Mommer, I., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above and belowground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 164–175.

Line105. How heating cables were buried? I hope sunny days were long enough to charge batteries and the batteries never failed

A detailed description of the experimental set-up and burying of the heating cables, including pictures, is provided in the reference indicated in L101 (Wang *et al.* 2017, including supplementary materials). The heating cables were buried into the soil from trenches excavated at two opposing sides of the experimental plots to minimize disturbance within the plots (i.e. disturbance of roots and microbial activity). For the heated cable treatment, the cables were connected to two solar panels of 85 W each, which were connected in parallel. The solar panels were installed with an angle of 60° to capture 20 hours of sunlight per day during the summer. No battery was included in the circuit. Thereby, the solar energy directly enlarged the natural ground heat flux, allowing for diurnal and seasonal variation in solar intensity.

Wang, P., Limpens, J., Mommer, I., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above and belowground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 164–175.

Line 166-170. I am afraid the random effects of the plot was not included in lme

For this statistical analysis – effects of permafrost thaw and fertilization treatments on soil temperature and thaw depth of the experimental plots – we used the average of the permafrost thaw depth and temperature data per plot as response variable. As indicated in subsection 2.3 of the manuscript, data were collected in 2013 (only temperature) and 2014 (temperature and thaw depth). Because we aggregated the data per experimental plot, we removed the random term (plot) from our analysis, in order to avoid model-overfitting.

Line 350-368. Discussion tries to relate no matter what I am trying to understand why this experiment could not be completed in less than four years when the year seems not to have any specific function, for example, repeated measurements? I know the fertilizer was applied twice, the second time after two years – other than that do not know why four years are emphasized? Stem traits did not show response even after four years anyway.

Although repeated measurements were not used in this study, the experimental duration over several years (i.e. 4 years) was important for many reasons. By running the experiment for four years, we aimed at reducing potential disturbance effects by the initial experimental setup (i.e. soil disturbance due to introduction of heating cables). Moreover, we selected a duration of several years considering the characteristics of tundra plant species, i.e. low rates of resource acquisition, growth and tissue turnover (Chapin 1980). This consideration was especially important for the shrub growth rate study associated with the experiment (Iturrate-Garcia et al., 2017). In this case, the experimental duration covered different climate conditions in the control plots across years and multiple annual growth rings. This fact allowed us to get insights into the effects of climatic conditions on growth rate (unpublished results).

As indicated in the answer addressing Referee 2's comment (L120), the aim of this study is to test treatment effects on plant traits by comparing treatment plots with control plots after 4 years of experiment and not to document relative changes for individuals over the experimental period. Because of this aim and the destructive sampling of individuals needed for the current study, we did not have repeated measurements across the years. Furthermore, we sampled only at the end of the experiment to avoid disturbing the setup.

The lack of response of stem traits, indeed, might be explained by the relative short term of the experiment, as explained in Lines 307-309. Wood tissue turnover is slower than leaf tissue (Negrón-Juárez et al., 2015). Thus, stem traits might require even more time (> 4 years) to show responses.

Chapin, F.S. III (1980). The mineral nutrition of wild plants. *Annual Review of Ecology and Systematics*, 11, 233–260.

Iturrate-Garcia, M., Heijmans, M.M.P.D., Schweingruber, F.H., Maximov, T.C., Niklaus, P.A., Schaeppman-Strub, G. (2017). Shrub growth rate and bark responses to soil warming and nutrient addition – A dendroecological approach in a field experiment. *Dendrochronologia*, 45, 12–22.

Negrón-Juárez, R.I., Koven, C.D., Riley, W.J., Knox, R.G., Chambers, J.Q. (2015). Observed allocations of productivity and biomass, and turnover times in tropical forests are not accurately represented in CMIP5 Earth system models. *Environmental Research Letters*, 10, 064017.

I do not see tables S2 S3 and fig. 5 s1 mentioned in the text

The tables S2 –S3 and figure S1 mentioned in the text are included in the supplementary material: <https://www.biogeosciences-discuss.net/bg-2019-498/bg-2019-498-supplement.pdf>

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## Responses to Michael Klinge (Referee 2)

### General comments

In this work the authors present their results of bio-ecological investigation on tundra shrubs. They used an experimental setup for examining the change of physiological plant conditions induced by actual climate change. The basic assumption was that climate warming leads to enhanced permafrost thaw, which simultaneously will add nutrients to the system. Leaf and stem traits of four different shrubs species were statistically analysed. The manuscript is generally well structured, clearly written and substantially justified by literature. The results about the potential adjustments of plant growing strategies contribute new insights for future environmental development in the subarctic region.

We thank Michael Klinge for his time to review our manuscript and for providing constructive and relevant comments, which will improve our manuscript.

### Specific comments

There are two general obstacles in the experiment setup, which I propose to consider more for discussion and conclusion: A main result of the experiment was that no significant response of plant traits was found due to permafrost thaw, whereas significant plant-trait response to fertilization was proofed. The general assumption was that the nutrient supply will increase caused by enhanced mineralization and thawed soil due to climate warming. This means that solely the soil heating would already lead to an increase of nutrient supply during the experiment. I am wondering, why an effect of increased nutrient supply is not observed in the data for the solely heating part of the experiment. Parallel soil analyses would have underlined the presumed causal chains.

We agree on the expected increase of nutrient supply in heating plots through two potential mechanisms, namely enhanced mineralization (if soil temperature increases) and thawed soil material containing nutrients. Parallel soil analyses have been performed and are reported in Wang *et al.*, 2017 (supplementary material). These analyses based on buried resin bags showed no increase of exchangeable nutrients in the non-fertilized unheated and heated plots for nitrogen and phosphorus. In the fertilized plots an increase in the top soil (5cm) was found for nitrogen (4x) and phosphorus (5x), and only slightly (but significantly) increased at a depth of 25 cm for nitrogen, but not for phosphorus. Increasing the energy input into the soil might result in a very low increase of temperature under humid conditions (i.e. high soil thermal conductivity) close to the permafrost table, where the cables were buried, as the energy might be partitioned towards the thawing process, and not towards heating the soil. Hence, the increase in mineralization rate might be rather limited in the plots with heated cables due to limited increase in temperature.

We will add a corresponding short section to the discussion, as suggested by the reviewer.

Wang, P., Limpens, J., Mommer, I., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above and belowground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 164–175.

The fertilization part of the experiment represents an extraordinary nutrient input into the system, which are marginal under natural conditions, when a slight input of nitrified dust from anthropogenic sources, desiccated lakes and desertificated landscapes is taken into account. The heating cables are buried in a depth of 15 cm. This brings along a systematic problem for the study design when compared to expected environmental changes under natural conditions. Climate warming controls soil temperatures along air temperatures. Thus, soil heating begins at the top surface and temperature changes move downward with decreasing amplitude. The high content of organic material in subarctic topsoil has a specific influence on the thermal conductivity into the subsoil. During

summer, it may have an isolating effect, when it becomes dried-up; during winter, the thermal conductivity increases caused by soil moisture content.

The fertilization indeed represents a large nutrient input into the ecosystem, much larger than the atmospheric nutrient input. The goal of adding fertilizer was to mimic future increased nutrient availability in the soil resulting from increased mineralization of nutrients in the soil organic layer in future warmer soils. The nutrient input is perhaps extraordinary compared to the atmospheric input, which is extremely low in the Arctic, but less so compared to soil nutrient availability and other fertilization experiments in Arctic tundra.

The fertilizer was added in the form of slow-release tablets. As the release rate depends on soil temperature, which is low at our study site, we applied a rather high dose. This dose should be seen as the maximum, probably not all of the nutrients have been released. When we performed measurements in the plots two years after application, we found some intact tablets in dry moss, indicating not all of the nutrients added had become available.

As stated by the reviewer, soil moisture conditions along the vertical profile will impact the energy partitioning within the soil towards heating and permafrost thawing. As mentioned above, the aim of this experiment was to simulate permafrost thaw and not soil warming. As stated in Wang *et al.*, 2017, ‘The deep soil heating treatment increased June–July thawing depth without increasing soil temperatures in the upper organic soil layer, for the first time enabling us to separate the effects of increased thawing depth from the effects of surface soil environmental changes in the tundra.’

We will add a section in the introduction to make this more explicit.

Wang, P., Limpens, J., Mommer, I., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above and belowground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 164–175.

## Technical corrections

L25: Here you are talking of “all four species” but until now you didn’t introduce them in the abstract.

We will add this information to L21: “...in four shrub species (*Betula nana*, *Salix pulchra*, *Ledum palustre*, *Vaccinium vitis-idaea*), which were sampled in the experimental plots”.

L59: Please give detailed information about the distinct methods used in “several experiments and satellite imagery”

We will add more information about the distinct methods used in “several experiments and satellite imagery”.

“Several warming experiments (Elmendorf *et al.*, 2012), satellite imagery – i.e. AVHRR, MODIS and Landsat multi-decadal records of the normalized difference vegetation index (NDVI) (Myers-Smith *et al.*, 2011) – and repeat multi-decadal aerial photography (Tape *et al.*, 2012) have shown effects of recent climate warming on tundra vegetation growth, productivity and distribution, especially on shrubs (Myers-Smith *et al.*, 2015; Myers-Smith & Hik, 2018)”.

Elmendorf, S.C., Henry, G.H.R., Hollister, R.D., Björk, R.G., Bjorkman, A.D., Callaghan, T.V., ..., Wookey, P.A. (2012). Global assessment of experimental climate warming on tundra vegetation: heterogeneity over space and time. *Ecology Letters*, 15, 164–175.

Myers-Smith, I.H., Elmendorf, S.C., Beck, P.S.A., Wilmking, M., Hallinger, M., Blok, D., ..., Vellend, M. (2015). Climate sensitivity of shrub growth across the tundra biome. *Nature Climate Change*, 5, 887–891.

Myers-Smith, I.H., Forbes, B.C., Wilmking, M., Hallinger, M., Lantz, T., Blok, D., ..., Hik, D.S. (2011). Shrub expansion in tundra ecosystems: dynamics, impacts and research priorities. *Environmental Research Letters*, 6, 045509 (15 pp).

Myers-Smith, I.H. and Hik, D.S. (2018). Climate warming as a driver of tundra shrubline advance. *Journal of Ecology*, 106, 887–891.

Tape, K.D., Hallinger, M., Welker, J.M., Ruess, R.W. (2012). Landscape heterogeneity of shrub expansion in Arctic Alaska, *Ecosystems*, 15, 711–724.

L120: When did you select and cut the individuals? I think after 4 years at the end of the experiment. This should be mentioned here. In addition: Why not selecting the individuals already in the beginning of the experiment; to measure some initial parameters such as plant height and LA? Then you would be able to document relative changes for individuals over the period?

Yes, the individuals were selected and sampled at the end of the experiment (year 4 of the experiment (2014)). We will add a sentence with this information to subsection “2.4 Study species and sampling”.

The aim of this study was to test treatment effects on plant traits by comparing treatment plots with control plots after 4 years of experiment and not to document relative changes for individuals over the experimental period. Because of this reason, we did not select individuals nor measured individual traits, such as height and LA, at the beginning of the experiment. Furthermore, we focused on assessing treatment effects on groups of traits, instead of on individual traits, to identify changes in shrub strategies. Changes on the selected strategies might provide insight on shrub community shifts in Arctic tundra under future climatic conditions expected for the Arctic, such as rising temperature and increasing permafrost thaw (IPCC 2013, Turetsky *et al.* 2020, Voigt *et al.*, 2017).

IPCC: *Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report of the Intergovernmental Panel on Climate Change* (eds. TF Stocker, D Qin, G-K Plattner, M Tignor, SK Allen, J Boschung, A Nauels, Y Xia, V Bex, and PM Midgley), Cambridge, United Kingdom and New York, NY, USA.

Turetsky, M.R., Abbot, B.W., Jones, M.C., Anthony, K.W., Olefeldt, D., Schuur, E.A.G., Grosse, G., Kuhry, P., Hugelius, G., Koven, C., Lawrence, D.M., Gibson, C., Sannel, A.B.K., McGuire, A.D. (2020). Carbon release through abrupt permafrost thaw. *Nature Geoscience*, 13, 138–143.

Voigt, C., Marushchak, M.E., Lamprecht, R.E., Jackowicz-Korczyński, Lingren, A., Mastepanov, M., Granlund, L., Cristensen, T.R., Tahvanainen, T., Martikainen, P.J., Biasi, C. (2017). Increased nitrous oxide from Arctic peatlands after permafrost thaw. *PNAS*, 114 (24), 6238–6243.

L135: Space between 1 and cm2

OK.



Referee's comments in black  
Authors' responses in blue

### Responses to Michael O'Brien (Short Comment)

I found the manuscript clear and well written. I think it provides a nice contribution to climate change impacts in the arctic system. Here are a few line-by-line comments.

We thank Michael O'Brien for his time and constructive comments, which will notably improve the clarity of the manuscript.

L12: I would not use the word adapt as that implies a genetic response, which is not tested here.

We will use "change" instead of "adapt" to avoid misunderstandings with genetic responses.

L62: I find this reference to Violle odd. I would say performance traits (detailed in Violle et al. 2007).

OK.

L66: Remove 'however'

OK

L120: When were these sampled? At the beginning and end of the experiment? Only after the 4 years?

The individuals were sampled at the end of the experiment (year 4 of experiment (2014)). We will add a sentence with this information to subsection "2.4 Study species and sampling".

L168 & L174: The model is actually block, heat treatment, fert treatment and treatment interaction. Not solely the interaction. I think the description of the analysis could use a bit of revision for clarity. It is often written in a condensed way that makes it tough to follow. I would add a bit of text to improve clarity of the model descriptions, especially for the contrasts.

We will correct the description of the model and revise it for clarity.

L167-L169: "[...]. The fixed terms of the models were block (factor with five levels), permafrost thaw treatment (two levels: heating, no-heating), fertilization treatment (two levels: fertilization, no-fertilization) and treatment interaction. [...]"

L172-L184: " We also used linear mixed-effect models to test the treatment effect on plant traits. Height, LA, bark thickness, and xylem diameter were log-transformed prior the statistical analysis to meet assumptions of linearity. First, we analysed plant traits of the four species together and then plant traits of each functional type (PFT; deciduous and evergreen). In the species analysis, we modelled each plant trait as a function of block (a fixed factor with five levels), permafrost thaw treatment (a fixed factor with three levels), fertilization treatment (a fixed factor with two levels), species (a fixed factor with four levels) and the interaction between treatments and species. In addition to these fixed terms, we also considered the interaction between species and block, which was a term recognised in the course of the statistical analysis to take into account species-specific trait differences among blocks. The random terms of the model were plot (factor with 30 levels) and the interaction of plot and species. In the PFT analysis, we modelled plant traits as a function of block, permafrost thaw treatment, fertilization treatment, PFT (a fixed factor with two levels),



species and the interaction between treatments, PFT and species. The random terms were plot and the interaction between plot and species. In both cases, we assessed if the effects of the permafrost thaw treatment on plant traits were due to the disturbance of the buried cables or the treatment *per se*. For that purpose, we split the three-level permafrost thaw factor into two contrasts of one degree of freedom (df) each, i.e. cable presence (heated and unheated cables vs. no-cable) and heating (heated cables vs. unheated cable and no-cable). We used the first contrast to assess cable effects (heating followed by cable presence) and the second contrast to assess treatment effects (cable presence followed by heating). After running these models for species and PFTs, we found that plant traits were significantly different among species, even between species within the same PFT. Consequently, we analysed the four species separately to maintain ecological information. We fitted block, permafrost thaw treatment, fertilization treatment and the treatment interaction as fixed terms and plot as a random term".

L203: I am not sure 'no-growing' is correct, maybe 'dormant'

We use the term 'no-growing', which is widely used in Arctic and alpine research. Some examples of this use can be found in Lin *et al.* (2011), Parmentier *et al.* (2011), Rumpf *et al.* (2014), van der Molen *et al.* (2007), Wang *et al.* (2016).

Lin, X., Zhang, Z., Wang, S., Hu, Y., Guangping, X., Luo, C., Chang, X., Duan, J., Lin, Q., Xu, B., Wang, Y., Zhao, X. and Xie, Z. (2011). Response of ecosystem respiration to warming and grazing during the growing seasons in the alpine meadow on the Tibetan plateau. *Agricultural and Forest Meteorology*, 792-802.

Rumpf, S.B., Semenchuk, S.D., Cooper, E.J. (2014). Idiosyncratic responses of High Arctic plants to changing snow regimes. *PLoS ONE*, 9(2): eB6281.

van der Molen, M.K.; van Huissteden, J., Parmentier, F.J.W., Petrescu, A.M.R., Dolman, A.J. *et al.* (2007). The growing season greenhouse gas balance of a continental tundra site in the Indigirka lowlands, NE Siberia. *Biogeosciences*, European Geosciences Union, 4 (6), pp. 985-1003.

Parmentier, F.J.W., van der Molen, M.K., van Huissteden, J., Karsanaev, S.A., Kononov, A.V., Suzdalov, D.A., Maximov, T.C. and Dolman, A.J. (2011). Longer growing seasons do not increase net carbon uptake in the northeastern Siberian tundra. *Journal of Geophysical Research*, 116, G04013.

Wang, P., Mommer, L., van Ruijven, J., Berendse, F., Maximov, T.C. and Heijmans, M.M.P.D. (2015) Seasonal changes and vertical distribution of root standing biomass of graminoids and shrubs at a Siberian tundra site. *Plant Soil*, 407, 55-65.

L266-267: Shrubs are plants, so this is redundant.

We will rephrase the sentence "Plant growth in high-latitude ecosystems is highly nutrient-limited (Billings & Mooney 1968; ...)".

L269: Confusing. Try : "Shrubs were released from growth limitation via nutrient addition, which was evidenced by. . ." or "Nutrient addition released shrubs from growth limitation as evidenced by. . ."

We will change the sentence to "Nutrient addition released shrubs from growth limitations as evidenced by the plant trait changes we found, such as ..."

L289-291: Which species would outcompete shrubs in this system? Which species will shade them out?

In Arctic tundra, graminoid species - particularly grass species - are expected to shade and outcompete shrubs, as suggested by results of warming and fertilization experiments carried out on tundra sites (Dormann and Woodin 2002, Gough and Hobbie 2003). In the study area, the species that are expected to outcompete shrubs are *Calamagrostis*

*holmii* Lange and *Eriophorum vaginatum* L. (Wang *et al.* 2017). We will add a sentence to the discussion to answer both questions.

Dormann, C.F. and Woodin, S.J. (2002). Climate change in the Arctic: using plant functional types in a meta-analysis of field experiments. *Functional Ecology*, 16, 4–17.

Gough, L. and Hobbie, E. (2003). Responses of moist non-acidic arctic tundra to altered environment: productivity, biomass, and species richness. *Oikos*, 103, 204–216.

Wang, P., Limpens, J., Mommet, L., van Ruijven, J., Nauta, A.L., Berendse, F., Schaepman-Strub, G., Blok, D., Maximov, T.C., Heijmans, M.M.P.D. (2017). Above- and below-ground responses of four tundra plant functional types to deep soil heating and surface soil fertilization. *Journal of Ecology*, 105, 947–957.

L301: evergreen and deciduous are not species but PFT and if you mean species then use among not between.

The sentence will be corrected: "...in the leaf economics spectrum both between PFTs, i.e. from evergreen to deciduous ... and within species...".

L310: Did you test wood density in tissues grown before and after treatment? It seems like the sampling strategy would allow partitioning the inner and outer stem to see differences.

We did not test wood density in tissues grown before and after treatment. Because of the sampling protocol applied (individuals were selected at the end of experiment), identifying tissues growing before and after the treatments were difficult. However, in another study based on this experiment (Iturrate-Garcia *et al.* 2017), the treatment effects were tested on inner and outer stem variables (i.e. bark thickness, xylem diameter, bark investment, wood biomass). In addition, annual growth rings of the four years before the experiment and the four years of experiment were tested, resulting in a significant increase on the distance between rings (i.e. growth rate) on shrubs growing on fertilized plots.

Iturrate-Garcia, M., Heijmans, M.M.P.D., Schweingruber, F.H., Maximov, T.C., Niklaus, P.A., Schaepman-Strub, G. (2017). Shrub growth rate and bark responses to soil warming and nutrient addition – A dendroecological approach in a field experiment. *Dendrochronologia*, 45, 12-22.

L320: It is not clear regarding the point that stem and leaf trade-offs operate independently. Please revise for clarity.

We will rephrase this sentence and the previous one for clarity.

Line 320: “However, our results showed that coordination between stem-height PC1 and leaf PC1 was only significant for half of the species. For *Betula nana* and *Vaccinium vitis-idaea*, the significant relationship between both axes suggested that these species coordinated stem and leaf traits (e.g. conservative trade-off at stem and leaf levels), resulting in a whole-plant strategy. The lack of coordination between stem and leaf traits for *Salix pulchra* and *Ledum pallustre* suggests that, for certain species, functional trade-offs at stem and leaf levels may operate partly independent (Fortunel, Fine & Baraloto, 2012).”

L349: Remove ‘also’

OK

L357: Confusing sentence.

We will revise the sentences included in Lines 356-357 for clarity:

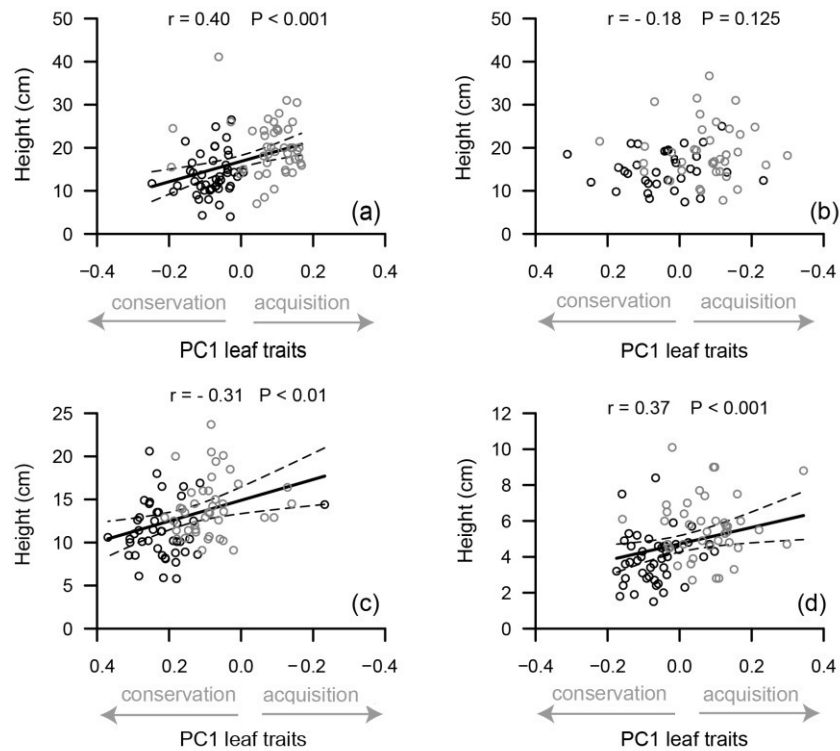
L356-357: “This depletion might result in reduced permafrost thaw through decreasing soil moisture, thermal conductivity, heat flux and temperature, which suggests that shrub shading might not be the only driver of the reduced permafrost thaw.”

L322-370: I found the discussion overly speculative in an effort to relate the trait responses into a climate-vegetation feedback. I would encourage the authors reduce the speculation or possibly present the information as potential scenarios of climate and vegetation responses.

This part of the discussion is meant to highlight how the results of this study based on detailed trait analyses and plant strategies support earlier findings that only covered part of the leaf and plant economic spectrum. We will revise the language of the indicated discussion section to address the reviewer’s comment.

Fig 5: Could you inverse two of the PC1 values so that the x-axis is always conservative on the left side and acquisitive on the right side? It would make it easier to read.

We will change Figure 5 to have the conservative strategy on the left side and acquisitive on the right side.



# Plant trait response of tundra shrubs to permafrost thaw and nutrient addition

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**Abstract.** Plant traits reflect growth strategies and trade-offs in response to environmental conditions. Because of climate warming, plant traits might ~~adapt~~change, altering ecosystem functions and vegetation–climate interactions. Despite important feedbacks of plant trait changes in tundra ecosystems with regional climate, with a key role for shrubs, information on responses of shrub functional traits is limited.

15 Here, we investigate the effects of experimentally increased permafrost thaw depth and (possibly thaw-associated) soil nutrient availability on plant functional traits and strategies of arctic shrubs in northeastern Siberia. We hypothesize that shrubs will generally shift their strategy from efficient conservation to faster acquisition of resources through adaptation of leaf and stem traits in a coordinated whole-plant fashion. Thereto we ran a 4-year permafrost thaw and nutrient fertilization experiment with a fully factorial block design and six treatment combinations – permafrost thaw (control, unheated cable, heated cable) × fertilization (no-nutrient addition, nutrient addition). We measured ten leaf and stem traits related to growth, 20 defence and the resource economics spectrum in four shrub species (*Betula nana*, *Salix pulchra*, *Ledum palustre* and *Vaccinium vitis-idaea*), which were sampled in the experimental plots. The plant trait data were statistically analysed using linear mixed-effect models and principal component analysis (PCA).

25 The response to increased permafrost thaw was not significant for most shrub traits. However, all shrubs responded to the fertilization treatment, despite decreased thaw depth and soil temperature in fertilized plots. Shrubs tended to grow taller, but did not increase their stem density or bark thickness. We found a similar coordinated trait response for all four species at leaf and plant level, i.e. they shifted from a conservative towards a more acquisitive resource economy strategy upon fertilization. In accordance, results point towards a lower investment into defence mechanisms, and hence increased shrub vulnerability to 30 herbivory and climate extremes.

Compared to biomass and height only, detailed data involving individual plant organ traits such as leaf area and nutrient contents or stem water content can contribute to a better mechanistic understanding of feedbacks between shrub growth strategies, permafrost thaw and carbon and energy fluxes. In combination with observational data, these experimental tundra

35 trait data allow for a more realistic representation of tundra shrubs in dynamic vegetation models and robust prediction of ecosystem functions and related climate-vegetation-permafrost feedbacks.

## 1 Introduction

Plants have different strategies to use resources to grow, reproduce, compete with neighbour plants and defend themselves against pathogens and herbivores (Bazzaz et al. 1987; Ordoñez et al. 2010). However, as resources – nutrients, water, and light – can have limited availability, plants are subject to trade-offs so that they have to allocate the resources to one function versus another (Grime 1977; Westoby et al. 2002; Reich 2014). Environmental changes, such as the ones promoted by climate warming (e.g. increasing amount of resources available in the soil), may modify these trade-offs and plant strategies (Grime 2006; Ordoñez et al. 2010).

Plant strategies and trade-offs can be identified by measuring plant traits and their correlations (Grime et al. 1997; Westoby et al. 2002). The so-called leaf economics spectrum is an example of how leaf traits show similar covariation across species according to the resource acquisition strategy, which goes from slow (i.e. resource conservation) to rapid resource acquisition (Díaz et al. 2016). Plant traits also determine plant responses to environmental factors and underpin their effects on ecosystem processes and services (Lavorel & Garnier 2002; Kattge et al. 2011; Soudzilovskaia et al. 2013). Analysing plant trait responses to climate warming can provide insight into future ecosystem structure and functioning (Díaz et al. 2007).

50 In low-resource environments such as the Arctic tundra, plants adopt a conservative strategy with low rates of resource acquisition, growth and tissue turnover (Chapin 1980). Low stature, small leaves of low specific leaf area (thick leaves with dense tissue) and long leaf life span reflect that conservative strategy (Reich, Walters & Ellsworth 1997; Cornelissen 1999). This strategy allows plants to allocate resources to other processes, such as defence against pathogens and herbivores, and confer stress resistance (Chapin, Autumn & Pugnaire 1993). Plants however might adopt a more acquisitive strategy under the environmental conditions projected for the Arctic in the course of this century (Post et al. 2019).

Surface air temperature in the Arctic has risen more rapidly than in other regions over the past decades and is projected to keep increasing; ( $\sim 3^{\circ}\text{C}$  by the year 2100 under emission scenario RCP4.5) (IPCC 2013). Ground heat flux and soil temperature are also expected to increase, accelerating permafrost thaw and soil organic matter mineralization (Rustad et al., 2001; Richter-Menge & Overland, 2010; Elmendorf et al., 2012). The release of nutrients trapped in the permafrost (Kokelj & Burn 2003; Weintraub & Schimel 2003; Schuur et al. 2009), together with an enhanced soil mineralization rate (Schmidt, Jonasson & Michelsen 1999; Walther et al. 2002), will increase nutrient availability for tundra vegetation (Keuper et al. 2012). Several warming experiments (Elmendorf et al. 2012), ~~and~~ satellite imagery – i.e. AVHRR, MODIS and Landsat multi-decadal records of the normalized difference vegetation index (NDVI) (Myers-Smith et al. 2011) – and repeat multi-decadal aerial photography (Tape et al. 2012) have shown effects of recent climate warming on tundra vegetation growth, productivity and distribution, especially on shrubs (Myers-Smith et al. 2011, 2015; ~~Elmendorf et al. 2012; Tape et al. 2012;~~

Myers-Smith & Hik 2018). Our current knowledge of tundra shrub responses to climate change concerns mainly their performance traits (detailed in sensu-Violle et al. (2007), especially plant height and biomass. ~~However, we~~ We still know precious little about the functional traits underpinning these responses or the effects expanding shrubs may have on ecosystem functions (but see Hudson, Henry & Cornwell 2011; Kremers, Hollister & Oberbauer 2015; Barrett & Hollister 70 2016). A recent pan-arctic plant trait sampling and analysis effort has revealed a generally strong spatial temperature-trait relationship, which was however mediated by soil moisture (Bjorkman et al., 2018a, Bjorkman et al. 2018b). This study also highlighted the limitations of the observational space-for-time substitution method and identified the need for experimental studies to elucidate intraspecific trait responses to environmental drivers. Shrub responses to climate may have consequences for the carbon cycle (e.g. increase carbon uptake) and the surface energy budget (e.g. decrease albedo), which in turn may 75 affect the regional climate (Eugster et al., 2000; Chapin, 2003; Beringer et al., 2005; Bonfils et al., 2012; Pearson et al., 2013; Juszak et al., 2017). A better understanding of shrub trait responses to climate, and shrub–climate interactions is fundamental to improve dynamic global vegetation models and predictions of vegetation shifts (Cramer et al. 2001; Doherty et al. 2010; Wullschleger et al. 2014).

The objective of ~~In~~-this study is, we to experimentally investigate the consequences of increased permafrost thaw and 80 nutrient addition on aboveground traits and trait coordination of tundra shrubs. We hypothesize that under simulated future environmental conditions (i.e. permafrost thaw and soil nutrient increase), (i) shrubs will shift their strategy from efficient conservation to faster acquisition of resources through adaptation of leaf and stem traits; and (ii) leaf traits, stem traits and height will show a coordinated response to these environmental changes as they all belong to the same overall resource economy dimension within the functional trait space. To test our hypotheses we ran a permafrost thaw and fertilization 85 experiment for four years in Siberia, and measured ten plant traits related to the leaf economics spectrum, growth and defence in tundra shrubs. Whereas most previous studies focused on one or two shrub species only, we explicitly compare the responses of four predominant species in order to find commonalities versus idiosyncrasies of intra- and interspecific trait response, as these are critical for upscaling from site level to tundra ecosystems at larger scales.

## 2 Materials and methods

### 90 2.1 Study area

The study area is located in the nature reserve of Kytalyk, in the continuous permafrost region of Yakutia, northeastern Siberia (70°49'N, 147°28'E, 10 m a.s.l.). Ice-rich permafrost and shallow active layers characterize the area (van Huissteden *et al.*, 2005; Iwahana *et al.*, 2014). The mean annual precipitation is 210 mm and the mean annual air temperature -13.1°C, with minimum and maximum monthly means of -33.5°C in January and 11.2°C in July (1980–2013, WMO station 21946, 95 Chokurdakh, monthly summaries of GHCN-D, NOAA National Climatic Data Center).

The experimental plots were placed on a moist acidic tussock tundra area, the soil of which is classified as Gelisol (Wang *et al.* 2017). In the Circumpolar Arctic Vegetation Map (Raynolds *et al.*, 2019) the vegetation type in this area is classified as tussock-sedge, dwarf-shrub, moss tundra. The main vegetation has a maximum canopy height of 25 cm and comprises sedge allies (mainly *Eriophorum vaginatum*), abundant deciduous and evergreen dwarf shrubs, bryophytes and lichens. The growing season lasts from the end of June to end of August in the study area (Parmentier *et al.* 2011). The slightly acidic soil (pH 6) has a silty-clay texture and high organic matter content (Blok *et al.* 2010, Bartholomeus *et al.*, 2012). The soil organic matter decomposition is low as indicated by the high average carbon to nitrogen ratio (22) and low cellulose to lignin ratio (2.4) (Iturrate-Garcia *et al.* 2016). At mid-growing season, the mean active layer thickness is 35 cm, increasing to about 50 cm at the end of the season.

## 105 2.2 Experimental design

To test whether climate change might have effects on shrub traits, we ran a permafrost thaw and nutrient fertilization experiment from 2011 to 2014 (Wang *et al.* 2017). The experiment had a fully factorial block design with five blocks, each with six plots of 1.5 × 1.5 m placed at randomly chosen locations in July 2010. Each block covered an area of approximately 10 m x 10 m. A picture of a typical block setup is contained in the supplementary material of Wang *et al.* 2017. Six treatment combinations – permafrost thaw (3 levels) × fertilization (2 levels) – were randomly assigned to the plots within blocks.

The permafrost thaw treatment consisted of no cable, unheated cable and heated cable. For this treatment, We we buried heating cables powered by solar panels at approximately 15 cm depth to increase the thaw depth in the unheated and heated plots in July 2010 (Wang *et al.* 2017). One year later, we connected the cables of the heated plots to solar panels in order to increase the thaw depth. ~~The permafrost thaw treatment consisted of no cable, unheated cable and heated cable.~~ The unheated cable plots served as a reference for the permafrost thaw treatment, while plots without cable were included to quantify possible disturbance effects of the cable alone. For the fertilization treatment (nutrient addition versus no addition), we applied slow-release NPK fertilizer tablets with micronutrients (Osmocote Exact Tablet, Scotts International, Heerlen, the Netherlands). The tablets were applied at approximately 5 cm depth at the start of the experiment (July 2011) and ~~two~~ years later again in 2013 (5.6 g N, 1.4 g P and 3.7 g K · m<sup>-2</sup> · yr<sup>-1</sup>), which increased the exchangeable nutrient content mainly in the upper soil layer (Wang *et al.*, 2017).

## 2.3 Soil temperature and thaw depth

Soil temperature of each plot was measured continuously in 2013-2014 at four depths (0 cm, 5 cm, 15 cm and 25 cm) using temperature loggers (iButton DS1922L/DS1921G, Maxim Integrated, USA). Thaw depth was measured twice in July 2014 by introducing vertically a metal rod with centimetre scale until hitting the frozen soil (Wang *et al.* 2017).



## 2.4 Study species and sampling

We investigated the response of four shrub species dominant at the study site and present in all experimental plots: two deciduous species *Betula nana* ssp. *exilis* (Sukazcev) Hultén and *Salix pulchra* Cham., and two evergreen species *Ledum palustre* ssp. *decumbens* (Aiton) Hultén and *Vaccinium vitis-idaea* L. (Fig. 1). The abundance of the four species was broadly similar in all plots, except for *S. pulchra*, which was less abundant.

At mid-growing season of the last year of experiment (31 July – 12 August 2014). We randomly selected six healthy-looking individuals (with less than 20% leaf damage) of each species per plot ~~at mid-growing season~~, except for *S. pulchra* for which only one to four individuals were present per plot. We cut the selected individuals 4 cm below the root collar after measuring their height. The sampling and transport of the plant samples followed the protocol for standardised trait measurements described in Pérez-Harguindeguy *et al.* (2013). Most of the plant traits were measured in the laboratory within a few hours.

## 2.5 Plant traits

We selected ten aboveground plant traits, which are related to and provide insight into shrub growth, defence and nutrient acquisition strategies, as well as into the interactions between tundra shrubs and carbon and energy fluxes. We measured the selected leaf and stem traits in each individual of the four shrub species one single time ~~(i.e. growing season of the last year)~~ based on sampled shrub individuals, see section 2.4 of experiment.

### 2.5.1 Height

Plant height was measured in the field as the vertical distance from the ground to the tallest vegetative tissue of the selected individuals (maximum vegetative height).

### 2.5.2 Leaf area (LA) and specific leaf area (SLA)

We cut two leaves per individual, including the petiole, from the top and bottom canopy layers. We scanned the leaves with a flatbed scanner (LiDE 70 Canon Inc., Japan, 300 dpi image resolution) calibrated with a 1 cm<sup>2</sup> reference. Then, we estimated LA by counting pixels using the software MatLab R2014a (The MathWorks, Inc., MA, USA). We oven-dried the scanned leaves (60°C, 72 h) and weighed them to determine SLA by dividing the LA of each leaf by its dry weight.

### 2.5.3 Leaf dry matter content (LDMC)

We followed a variation of the partial rehydration method to determine LDMC using the same leaves as for LA (Vendramini *et al.* 2002; Vaieretti *et al.* 2007). To assure maximum hydration, we cut whole individuals in the morning, wrapped the samples in moist paper and put them in sealed plastic bags (Pérez-Harguindeguy *et al.* 2013). We kept the samples in the

dark at low temperatures until they were weighed within the following six hours to obtain fresh mass. The individual leaves  
155 were re-weighed after oven-drying them (60°C, 72 h). LDMC was the dry mass of a leaf divided by its fresh mass.

#### **2.5.4 Leaf nitrogen concentration (LNC)**

Oven-dried leaves were milled and leaf carbon and nitrogen concentrations determined by dry combustion (TruSpec Micro-  
CHN analyser, Leco Corporation, MI, USA) in samples of 2 mg. Then, the carbon to nitrogen ratio (C:N) was calculated.

#### **2.5.5 Leaf phosphorus concentration (LPC)**

160 We used a colorimetric assay employing ammonium heptamolybdate to determine LPC. Milled samples of 0.05 g were  
combusted in a muffle furnace (B180 Nabertherm, Germany) programmed with one-hour heating up ramp to 600°C and two  
hours and a half at 600°C. We added 2 ml of 0.1 M H<sub>2</sub>SO<sub>4</sub> to the ashes, followed by 5 ml of distilled water, and filtered the  
suspension (Macherey Nagel MN615). The phosphorus in the extracts was determined using a continuous flow analyser  
(Skalar Analytical B.V., the Netherlands) calibrated with KH<sub>2</sub>PO<sub>4</sub> standards.

#### **165 2.5.6 Stem-specific density (SSD)**

We cut approximately 3 cm long sections of the main stem at one third of the stem length and removed the bark. We  
measured the diameter and length of the stem sections without bark, oven-dried (60°C, 72 h) and weighed them. SSD was  
determined by dividing the dry mass of a section by its volume.

#### **2.5.7 Stem water content (SWC)**

170 We weighed the sections used for SSD before and after oven-drying them. SWC was estimated as the difference between  
fresh and dry weight divided by the dry weight.

#### **2.5.8 Xylem diameter and bark thickness**

Samples including the 2 cm above and below the root collar of the main stem were taken and preserved in ethanol (40% vol.  
aqueous solution) until laboratory processing. We cut thin sections of 20 – 30 µm along the root collar of each individual and  
175 placed them on microscope slides. We photographed and measured xylem diameter and bark thickness following the  
protocol described in Iturrate-Garcia *et al.* (2017).

### **2.6 Statistical analysis**

To test if soil temperature and thaw depth were affected by permafrost thaw and fertilization treatments, we used linear  
mixed-effect models fitted in asreml (ASReML 3.0, VSN International Ltd., UK). The fixed terms of the models were block  
180 (factor with five levels), ~~and the interaction among~~ permafrost thaw treatment (two levels: heating, no-heating), ~~and~~

fertilization treatment (two levels) and treatment interaction. For the analysis, we averaged the thaw depth values per plot. The soil temperature values were averaged by growing and no-growing season per plot and depth class. Because we aggregated the data per experimental plot, we removed the random term (plot) from the analysis, in order to avoid model-overfitting.

185 We also used linear mixed-effect models to test the treatment effect on plant traits. Height, LA, bark thickness, and xylem diameter were log-transformed prior the statistical analysis to meet assumptions of linearity. First, we analysed plant traits of the four species together and then plant trait of each functional type (PFT; deciduous and evergreen). In the species analysis, we modelled each plant trait as a function of ~~The fixed terms of the models were~~ block (a fixed factor with five levels), ~~the interaction among~~ permafrost thaw treatment (fixed factor with three levels), fertilization treatment (fixed factor with two levels), ~~and~~ species (fixed factor with four levels; first analysis) and the interaction between treatments and species. ~~or PFT (two levels) and species (PFT analysis).~~ For the first analysis, we also considered ~~In addition to these fixed terms, we also considered~~ the interaction between species and block, which was a ~~(term recognised in the course of the statistical analysis to take into account species-specific trait differences among blocks).~~ The random terms of the model were plot (factor with 30 levels) and the interaction of plot and species. In the PFT analysis, we modelled plant traits as a function of block, permafrost thaw treatment, fertilization treatment, PFT (a fixed factor with two levels), species and the interaction between plot and species. In both cases, we ~~tested for~~ assessed if the effects of the permafrost thaw treatment on plant traits were due to the disturbance of the buried cables or the treatment per se. ~~For that purpose, we split~~ of increased thaw depth and cable disturbance by splitting the three-level permafrost thaw factor into two contrasts of one degree of freedom (df) each, i.e. ~~presence of cable~~ presence (heated and unheated cables vs. no-cable) and heating (heating cables vs. unheated cable and no-cable). We used the first contrast to assess Cable disturbance effects ~~were tested by fitting~~ (heating (heated cable vs. unheated cable and no cable) followed by cable presence) (heated and unheated cable vs no cable), whereas increased and the second contrast to assess thaw depth treatment effects ~~were tested by fitting~~ (cable presence followed by heating). After running these models for species and PFTs, w~~We~~ found that plant traits were significantly different among species, even between species within the same PFT. Consequently, we analysed the four species separately to maintain ecological information. In this case, w~~we~~ fitted block, permafrost thaw treatment, fertilization treatment and the interaction between the permafrost thaw and fertilization treatments as fixed terms and plot as a random term.

200 In order to explore shrub plant strategy and its change with treatments, standardized (Z-scored) plant trait data were subjected to a principal component analysis (PCA; vegan package version 2.4-0; Oksanen 2016). We only considered the fertilization treatment (nutrient addition and no addition) in the PCA, as most traits were not responsive to the permafrost thaw treatment (see Results). We performed a separate analysis for leaf traits (SLA, LDMC, LNC, LPC and C:N), and one for stem traits (SSD, bark thickness, xylem diameter and SWC) and height. Scores and variable loadings resulting from the PCA were scaled for data visual depiction.

To test for relationships between leaf economics, and stem traits and height, we used linear mixed-models. We extracted the loadings of the first principal component axes (PC1) of the leaf trait and the stem trait-height PCA. The response variables in

215 our models were height and stem trait-height PC1 loadings. Block and leaf trait PC1 loadings were set as fixed terms and plot as random effect. The significance of the linear relationships between variables was analysed using Pearson's correlation coefficients in addition to the linear mixed-effect models.

All data were analysed using R 3.4.1 (<http://r-project.org>).

### 3 Results

#### 220 3.1 Treatment effects on soil temperature and thaw depth

Permafrost thaw and fertilization treatments affected soil temperature and thaw depth (Table 1). The soil temperature during the growing season was significantly higher in heated plots than in unheated plots. The soil temperature at 5 cm depth was 0.6°C higher ( $F_{1,29} = 9.00$ ,  $p < 0.05$ ), at 15 cm 1.1°C ( $F_{1,29} = 17.97$ ,  $p < 0.001$ ) and at 25 cm 0.8°C ( $F_{1,29} = 14.02$ ,  $p < 0.01$ ). The difference of surface soil temperature (0 cm) between heated and unheated plots was not significant. During the no-  
225 growing season, temperature differences were significant at all the depths. The soil temperature at 0 cm was 1.4°C higher in heated plots than in unheated plots ( $F_{1,29} = 16.2$ ,  $p < 0.01$ ), at 5 cm 1.3°C ( $F_{1,29} = 26.1$ ,  $p < 0.001$ ), at 15 cm 1.3°C ( $F_{1,29} = 17.5$ ,  $p < 0.01$ ) and at 25 cm 1.2°C ( $F_{1,29} = 16.9$ ,  $p < 0.01$ ). The difference in soil temperature between fertilized and unfertilized plots was also significant at all the depths, but only during the growing season. The soil temperature was lower in the fertilized plots: at 0 cm 0.9°C lower ( $F_{1,29} = 11.6$ ,  $p < 0.01$ ), at 5 cm 1.0°C ( $F_{1,29} = 19.1$ ,  $p < 0.01$ ), at 15 cm 0.4°C ( $F_{1,29}$   
230 = 12.7,  $p < 0.01$ ) and at 25 cm 0.5°C ( $F_{1,29} = 6.24$ ,  $p < 0.05$ ). The thaw depth was 10.7 cm deeper in heated plots than in unheated plots ( $F_{1,29} = 24.6$ ,  $p < 0.001$ ), but 3.9 cm shallower in fertilized plots than in unfertilized plots ( $F_{1,29} = 5.40$ ,  $p < 0.05$ ). Fertilization treatment effects on soil temperature and thaw depth did not depend on the permafrost thaw treatment.

#### 3.2 Treatment effects on leaf and stem traits and plant height

The permafrost thaw treatment had no significant effect on most shrub traits. Only LA responded significantly to the  
235 permafrost thaw treatment ( $F_{1,28} = 18$ ,  $p < 0.001$ ) when analyzing all four species together. At the species level, the permafrost thaw treatment affected only LA of *S. pulchra* and *L. palustre*. Individuals of both species had greater LA on heated plots than on control and unheated plots (Table 2). The permafrost thaw treatment only increased SWC for *S. pulchra* ( $F_{1,28} = 12.8$ ,  $p < 0.01$ ). Neither the effect of the combination of treatments (permafrost thaw × fertilization) nor the disturbance caused by the buried cables were significant for most measured leaf and stem traits. Exceptions were a  
240 significant treatment combination effect on bark thickness of *B. nana* ( $F_{1,25} = 4.54$ ,  $p < 0.05$ ) and *L. palustre* ( $F_{1,25} = 8.15$ ,  $p < 0.01$ ), and LA of *S. pulchra* being negatively affected by the buried cables (Table 2).

The fertilization treatment had a significant effect on all leaf traits, height, and SWC, but not on bark thickness, xylem diameter or SSD, when the four species were analysed together (results not shown). At the PFT level, traits were significantly different between deciduous and evergreen species, except for LDMC and SSD. We also found that the  
245 fertilization effects on LA, LNC, LPC, C:N and SWC differed between PFTs (Table S2). The relative increase of LA and

decrease of C:N with fertilization was greater for evergreen than for deciduous species. For LNC, LPC, and SWC, the increase was greater for deciduous than for evergreen species. At the species level, the fertilization effect on LA, LNC and C:N was significant for all four shrub species (Table 2). Fertilization effects were also significant for SLA, LPC and LDMC of all species except for *S. pulchra* (Table 2). Leaves in the fertilized plots were bigger and thinner (higher SLA), had higher nutrient concentration (LNC, LPC) and lower LDMC and C:N than leaves in unfertilized plots (Table 2). For stem traits, the fertilization treatment significantly increased the SSD of *B. nana* ( $F_{1,29} = 10.1$ ,  $p < 0.01$ ) and SWC of both deciduous species (*B. nana*:  $F_{1,29} = 17.8$ ,  $p < 0.001$ ; *S. pulchra*:  $F_{1,29} = 13.9$ ,  $p < 0.01$ ). Xylem diameter and bark thickness responses to nutrient addition were not significant.

### 3.3 Coordinated trait response to fertilization

In the leaf trait PCA with all four species combined, shrub individuals were separated into species with low overlap along the first principal component axis (PC1) (Fig. 2). PC1 explained 64% of the variation among individuals and was mainly related to leaf nutrient content (LNC, LPC) and C:N. We found *B. nana* at the lower end of PC1, associated with high SLA and leaf nutrient concentrations, and *V. vitis-idaea* at the upper end of the axis. *B. nana* was the species with the widest range along PC1. The second PC axis (PC2) explained 19% of the variation and was mainly related to LDMC. Under nutrient addition, we observed a similar trait change for all four species. Leaves on fertilized plots had lower LDMC and C:N and higher LNC, LPC and SLA than leaves on unfertilized plots (Fig. 2).

Similar leaf trait space occupation was found when we ran the PCA for each species separately (Fig. 3). PC1 explained a slightly greater amount of total variance among individuals for the evergreen species (*L. palustre* 65% and *V. vitis-idaea* 60%) than for the deciduous species (*B. nana* 54% and *S. pulchra* 41%). Individuals were separated into two clusters along PC1 corresponding to individuals from fertilized and unfertilized plots. PC2 explained 17% and 18% of the variation among individuals for *L. palustre* and *V. vitis-idaea*, respectively, and 20% for deciduous species. The main results were maintained when we excluded LNC from above analysis, showing that PC1 was not driven by the potential correlation of C:N and LNC (Table S3).

Similarly, we ran a PCA for stem traits and plant height for each of the four species (Fig. 4). For these traits, individuals overlapped more on the PCA ordination plane. However, there was a trend towards taller individuals having lower SSD and higher SWC in the fertilized plots for three species, but not for *V. vitis-idaea*. Indeed the stem trait-height space was generally similar for all the species except *V. vitis-idaea*. PC1 explained slightly more variation among individuals than PC2, especially for deciduous species.

### 3.4 Plant strategies – correlation of leaf traits with stem traits and plant height

We found significant correlation between PC1 of the leaf trait PCA (leaf PC1) and plant height for all species, except for *S. pulchra* (Fig. 5). We also found a significant correlation between leaf PC1 and stem trait-height PC1 for *B. nana* and *V. vitis-*

*idaea* (Fig. S1). Individuals found in the upper range of the stem trait-height PC1 (high values for height, xylem diameter and bark thickness) were also found on the upper extreme of leaf PC1 (high values of LNC and LPC).

## 4 Discussion

280 We experimentally tested the effects of increased thaw depth and nutrient availability on plant traits of four tundra shrub species. While no strong responses to permafrost thaw were observed, our findings did show a coordinated response of leaf traits to fertilization, i.e. from a strategy of conservation of resources towards more rapid resource acquisition at leaf level, as we had hypothesised. Stem traits also tended towards a coordinated response to fertilization, though to a lesser extent. Moreover, one of the two deciduous (i.e., *Betula nana*) and one of the two evergreen species (i.e., *Vaccinium vitis-idaea*)  
285 showed a coordinated response of leaf and stem traits to fertilization along the same resource economics axis.

### 4.1 Treatment effects on plant traits

We expected that permafrost thaw and fertilization treatments would affect plant traits. However, our results showed that most of the plant traits responded only to the shallow nutrient addition. Plant growth in high-latitude ecosystems is highly nutrient-limited ~~and so is shrub growth~~ (Billings & Mooney 1968; Shaver & Chapin 1980; Epstein *et al.* 2000). Nutrient  
290 addition releases shrubs from this limitation and promotes their growth and biomass production (Chapin & Shaver 1996; DeMarco *et al.* 2014; Iturrate-Garcia *et al.* 2017). ~~Nutrient addition released The shrubs from growth limitation release by adding nutrients is reflected in the~~ as evidenced by the plant trait changes we found, such as greater height, SLA and leaf nutrient concentration (Hudson, Henry & Cornwell 2011; Reich 2014). It is notable that in this short-term, ~~strong fertilization treatment~~ reduced soil temperatures and permafrost ~~thaw~~ depth were measured under the strong fertilization  
295 treatment. Despite less favourable soil physical conditions, shrubs followed a more acquisitive growth strategy under fertilization as compared to the permafrost thaw treatment which had higher soil temperatures and deeper thaw depth. The fact that plant traits were less responsive to permafrost thaw than to fertilization might be explained by the relatively large amount of nutrients added to the fertilized plots ~~with the fertilization treatment~~. Parallel soil analyses were performed in the experimental plots by Wang et al. (2017; supplementary material). These analyses, based on buried resin bags (Giblin et al. 1991), showed no increase of exchangeable nutrients in the non-fertilized unheated and heated plots for nitrogen (N) and phosphorus (P) at any of the analysed depths. In the fertilized plots, they found that the nutrient amount in the top soil layer (< 5 cm) increased by four (N) and five (P) times. A significant but smaller increase was found only for N at deeper layers (25 cm). These findings, therefore, show that the nutrient amount added to the plots with the fertilization treatment was greater than the nutrient amount released by the thawing treatment.  
300 In plots with heated cables, enhanced nutrient availability is expected through acceleration of soil organic matter mineralization ((Knorr et al. 2005; Schaeffer et al. 2013), besides permafrost thaw and related release of nutrients. This amount may be greater than the nutrient release by permafrost thaw (Giblin et al. 1991; Hartley et al. 1999; Schaeffer et al.  
305

2013). Furthermore, the depth of soil layers at which nutrients were available for plants and of shrub rooting might also explain the different trait responses to the treatments. Most of the root biomass of the shrub species studied occurs at shallow soil depth (ca. 5–10 cm), which is shallower than the permafrost thaw depth during the growing season (Churchland *et al.* 2010; Keuper *et al.* 2012; Wang *et al.* 2017). In addition to permafrost thaw and related release of nutrients, soil warming might also enhance the nutrient availability through acceleration of soil organic matter mineralization (Knorr *et al.* 2005). Hartley *et al.* (1999) found effects of soil warming on subarctic shrub growth by using heating cables buried at 5 cm depth, which increased the soil temperature by 5°C. In our study, however, the heating cables were buried at 15 cm below the surface in order to increase the permafrost thaw without increasing the soil temperature of the shallower layers (< 15 cm). Thereby, most warming was in the mineral soil layers below 15 cm, whereas the increase of soil temperature in the shrub root layer was lower than the threshold (1°C or greater) needed for increasing nutrient mineralization (Schmidt *et al.* 1999). In addition, soil moisture conditions along the vertical profile affect the energy partitioning within the soil. Humid conditions close to the permafrost table, where the cables were buried, can promote energy partitioning towards permafrost thawing instead of towards soil heating. Under these conditions, increasing the input energy of the soil might result in a low temperature rise (i.e. high soil thermal conductivity), limiting in turn greater mineralization rates in plots with heated cables. Furthermore, the depth of soil layers at which nutrients were available for plants and of shrub rooting might also explain the different trait responses to the treatments. Most of the root biomass of the shrub species studied occurs at shallow soil depth (ca. 5–10 cm), which is shallower than the permafrost thaw depth during the growing season (Churchland *et al.* 2010; Keuper *et al.* 2012; Wang *et al.* 2017).

#### 4.2 Coordinated leaf trait response to nutrient addition

Resource availability is thought to be one of the main drivers of plant strategy selection (Grime 2006; Ordoñez *et al.* 2010). In arctic tundra, where resource availability is low, shrub species adopt a conservative strategy with slow growth and tissue turnover, which enhances plant survival under harsh conditions (Chapin *et al.* 1993). However, the “slow traits” associated with the conservative strategy are disadvantageous in case of higher resource availability as shrub species could be outcompeted (e.g. through shading) by other species with faster growth and biomass production (Reich 2014). In Arctic tundra, graminoid species – particularly grass species – are expected to shade and outcompete shrubs, as suggested by warming and fertilization experiments carried out at tundra sites (Dormann and Woodin 2002; Gough and Hobbie 2003; Wang *et al.* 2017). Our results show that species with similar resource economic strategies cluster into groups – deciduous and evergreen plant functional types – defined by their covarying leaf traits (Reich *et al.* 1997, 1999). On unfertilized plots, the deciduous shrub species *B. nana* and *S. pulchra* were characterized by leaf traits associated with faster resource acquisition: high SLA and leaf nutrient concentration and low LDMC and C:N. In contrast, the evergreen shrub species *L. palustre* and *V. vitis-idaea* were characterized by leaf traits associated with resource conservation, as expected due to a slower tissue turnover as compared to deciduous shrubs (Chapin & Shaver 1996).



We found different plant trait responses to fertilization with PFT for most leaf traits. Despite these differences, the increase of nutrients promoted a common coordinated response of leaf traits of all species, which reflects a change in resource economics from conservation to faster acquisition, even in the case of the evergreen species. Thus, there appears to be a comparable shift towards resource acquisitiveness in the leaf economics spectrum both between ~~species~~PFTs, i.e. from evergreen to deciduous (Wright *et al.* 2004; Freschet *et al.* 2010; Díaz *et al.* 2016), and within species (this study, Aerts *et al.* 2012). Since deciduous shrubs have been found to expand much more than evergreen shrubs in biomass and abundance in response to fertilization, both in Eurasian and North American tundra (van Wijk *et al.* 2003), our findings point to a possible important positive feedback between species turnover and intraspecific change with respect to resource economics traits.

#### 4.3 Stem traits response to nutrient addition

350 Stem traits were less responsive to treatments than leaf traits, which might be explained by the relative short term of the experiment. Turnover of wood tissue is slower than of leaf tissue (Negrón-Juárez *et al.* 2015). Thus, stem traits might require more time to show responses. Furthermore, the age heterogeneity of the selected shrubs might mask stem trait responses. Older individuals have higher SSD and bark thickness than younger ones (Woodcock & Shier 2002; Patiño *et al.* 2009; Poorter *et al.* 2014). Therefore, stem trait responses might become statistically significant when longer-term experiments are  
355 run and shrubs within the same age class (i.e. similar stem diameter) are selected.

Under nutrient addition, we found that coordinated stem trait response tended towards greater height and SWC and lower SSD. These findings are in line with previous studies showing a negative relationship between wood density and water content (Dias & Marengo 2014). Stems with lower SSD have less space filled with cell walls than those with higher SSD and therefore more water can be stored within the stem wood (McCulloh *et al.* 2011; Dias & Marengo 2014). Woody species  
360 with denser wood grow slower, have less wood water content and produce smaller and thicker leaves, which might be associated with a whole-plant strategy (Bucci *et al.* 2004; Wright *et al.* 2004; Ishida *et al.* 2008; Chave *et al.* 2009). However, our results showed that coordination between stem-height PC1 and leaf PC1 ~~\_, which reflect different trade-offs, were-was~~ only significant for half of the species. For *Betula nana* and *Vaccinium vitis-idaea*, the significant relationship between both axes suggests that these species coordinated stem and leaf traits (e.g. conservative trade-off at stem and leaf  
365 levels), resulting in a whole plant strategy. The lack of coordination between stem and leaf traits for *Salix pulchra* and *Ledum pallustre* suggests that, for certain species, functional trade-offs At stem and leaf levels, ~~functional trade-offs~~ may operate partly independently (Fortunel, Fine & Baraloto 2012).

#### 4.4 To grow or to defend

Our findings suggest that shrubs will grow taller, acquire more resources and allocate them to produce larger leaves at lower  
370 cost (thinner leaves with lower LDMC and C:N). These changes in plant traits, together with an expected faster growth, will come with a cost for shrubs: a decrease of their stress resistance (growth-defence trade-off) (Chapin *et al.* 1993; Chave *et al.* 2009; Iturrate-Garcia *et al.* 2017). The faster resource acquisition will make shrubs more vulnerable to herbivory due to

375 higher leaf nitrogen content (Mattson 1980; Díaz *et al.* 2016) and to adverse environmental conditions (i.e. low nutrient availability) as consequence of low nutrient tissue reserves (Reich 2014). We also found that shrubs with more rapid resource acquisition grew taller but without increasing their bark thickness and SSD, which might enhance shrub vulnerability to pests, mechanical and hydraulic failure, and extreme climatic events (Baraloto *et al.* 2010; Reich 2014; Díaz *et al.* 2016).

#### 4.5 Shrub–climate feedbacks

380 Vegetation is strongly coupled with environmental conditions (Wookey *et al.* 2009; Medinski *et al.* 2010). Shrubs will be affected by climate warming, with resultant changes in plant strategy and traits, affecting species diversity and ecosystem functions, such as carbon cycling and the surface radiation budget (Chapin *et al.* 1996; Beringer *et al.* 2005; Myers-Smith *et al.* 2011). The carbon uptake associated with increasing shrub growth and biomass production together with longer turn-over time due to carbon storage in branches as compared to leaf material will affect the carbon cycle (Hobbie, Nadelhoffer & Högberg 2002; Mack *et al.* 2004). Moreover, shrub trait changes may as well affect ecosystem processes. The production of low-cost tissues might accelerate litter decomposition because these tissues are easier to decompose than expensive ones (McLaren *et al.* 2017).

Our results suggest that tundra shrubs will be affected by increased nutrient availability in shallow soil layers. Deeper-rooting species, such as graminoids, may benefit more from nutrient release by permafrost thaw in deep soil layers (Keuper *et al.* 2017; Wang *et al.* 2017). In competition with graminoids, shrubs will pre-empt nutrient and light resources by growing faster and taller, producing denser canopies and leaves with greater photosynthetic area (Chapin & Shaver 1996; Hudson *et al.* 2011; Elmendorf *et al.* 2012; Díaz *et al.* 2016). Bryophyte and lichen diversity is expected to decline due to the increase of shading and litter deposition associated with those changes (Cornelissen *et al.* 2001; van Wijk *et al.* 2003; Elmendorf *et al.* 2012; Lang *et al.* 2012). As a consequence of the cryptogam decrease, the thermal insulation of the permafrost might be reduced (Blok *et al.*, 2011a), promoting permafrost thaw and the release of carbon (e.g. in the form of methane) to the atmosphere (Schuur *et al.* 2008; Schaefer *et al.* 2011). However, ~~also~~ shrub cover increase has been reported to reduce summer permafrost thaw locally (Blok *et al.*, 2011b; Nauta *et al.*, 2015; Wang *et al.*, 2017). While these studies discussed shading effects as main cause, our detailed trait analysis suggests additional mechanisms associated with water demand. Shrubs under nutrient addition showed greater SLA, lower LDMC and higher water content of leaves and stems pointing towards enhanced water demand through higher photosynthetic potential and evapotranspiration. The higher water demand might deplete soil water resources, as suggested by the lower soil moisture and summer soil temperature in the fertilized plots (for detailed results see supplementary material in Wang *et al.*, 2017), where deciduous shrubs increased most. This depletion might result in reduced permafrost thaw through decreasing soil moisture, heat-thermal conductivity, heat flux and temperature, which suggest that shrub shading might ~~and in turn reduced permafrost thaw. Shading by increased shrub cover might therefore~~ not be the only driver of the reduced permafrost thaw. Water demand by plants, especially shrubs, might be at least as important, as also documented in Juszak *et al.*, 2016. Interestingly, soil moisture has been found as a potential

growth co-limiting factor of tundra shrubs (Blok *et al.*, 2010, Myers-Smith *et al.*, 2015). However, shrubs might be released from water limitation by the predicted concomitant increase in precipitation. Thereby, related effects on shrub growth, community composition and feedbacks with the permafrost system and the atmosphere remain to be tested.

## 5 Conclusions

410 The climatic conditions projected for the Arctic, the shrub growth sensitivity to climate, and the importance of shrub–climate feedbacks for ecosystem functioning suggest that a special effort should be made to better understand future tundra changes and adaptation to the new climatic conditions. Here, we presented the response of a wide set of traits of selected dominant species in tussock tundra to permafrost thaw and increased nutrient availability. This response can be considered a step towards more realistic dynamic global vegetation models, although generalization should be considered cautiously due to the  
415 short term of the response, the spatial heterogeneity of Arctic regions and the complexity of shrub–climate feedbacks. According to our results, coordinated trait responses representing the whole plant (including wood and bark traits as in our study and ideally also root traits) instead of single trait responses are needed for a more robust prediction of shifts in vegetation, ecosystem processes and related climate–vegetation feedbacks.

## Data availability

420 All data presented in this paper will be available in the DryAd repository.

## Author contributions

MI and GS conceived the idea and methods of the study; GS and PAN obtained the grant that funded this research. MMPDH conceived the experimental design; MI collected the data; FHS instructed and contributed to the dendroecological work; MI and PAN analysed the data; MI led the writing of the manuscript; GS, PAN, MMPDH and JHCC contributed critically to the  
425 drafts. All authors gave final approval for publication.

## Competing interests

The authors declare that they have no conflict of interest.

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**Table 1: Average soil temperature (standard deviation) and average thaw depth (standard deviation) of the five experimental blocks for plots grouped into no-fertilization (NF) and fertilization treatments (F). Data are grouped by growing season (T<sub>Jun-Aug</sub>) and no-growing season (T<sub>Sep-May</sub>) and depth (0, 5, 15 and 25 cm).**

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	NH	H	NF	F
T <sub>Jun-Aug</sub> (°C)				
0 cm	8.4 (1.9)	8.5 (2.2)	8.8 (2.1)	<b>7.9** (1.8)</b>
5 cm	4.7 (1.9)	<b>5.3* (2.0)</b>	5.4 (2.1)	<b>4.4** (1.8)</b>
15 cm	1.4 (1.4)	<b>2.5*** (1.7)</b>	2.2 (1.6)	<b>1.8** (1.4)</b>
25 cm	0.4 (1.2)	<b>1.2** (1.5)</b>	0.9 (1.5)	<b>0.4* (1.2)</b>
T <sub>Sep-May</sub> (°C)				
0 cm	-13.1 (7.2)	<b>-11.7** (7.0)</b>	-12.4 (7.2)	-12.7 (7.1)
5 cm	-11.8 (6.8)	<b>-10.5*** (6.7)</b>	-11.2 (6.9)	-11.4 (6.8)
15 cm	-11.0 (6.7)	<b>-9.7*** (6.6)</b>	-10.5 (6.7)	-10.6 (6.7)
25 cm	-10.6 (6.6)	<b>-9.4** (6.5)</b>	-10.1 (6.6)	-10.2 (6.6)
Thaw depth (cm)				
	37.0 (6.6)	<b>47.7*** (3.2)</b>	42.5 (7.6)	<b>38.6* (7.2)</b>

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\*P < 0.05 \*\* P < 0.01 \*\*\*P < 0.001

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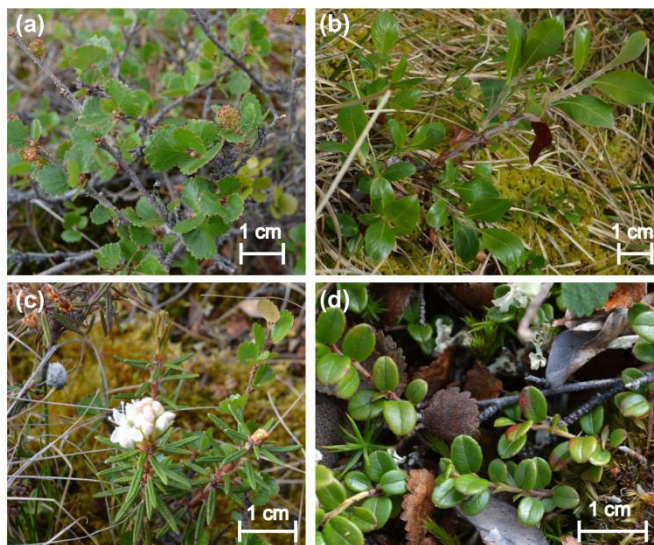
740 **Table 2: Effects of fertilization (Fert), cable disturbance (Ca) and soil heating (H) on leaf traits of each shrub species (*B. nana***  
**(Betn), *S. pulchra* (Salp), *L. palustre* (Ledp) and *V. vitis-idaea* (Vacv)). Treatment columns show the average and standard error of**  
**the response variables for no fertilization (NFert), fertilization (Fert), no heating cables (Ct), unheated cables (Ca) and heated**  
**cables (H). LMM columns show the Wald test outputs for our linear-mixed models (\*\*\* P < 0.001, \*\* P < 0.01 and \* P < 0.05).**  
745 **Significant effects are in bold. Treatment combination effect (heating × nutrient addition) was not included as it was significant**  
**only for LA of Betn (F<sub>1,25</sub> = 15.1 , P < 0.01).**

	Treatment					LMM		
	Fertilization		Permafrost thaw			Fert	Ca	H
	NFert	Fert	Ct	Ca	H	F <sub>1,29</sub>	F <sub>1,28</sub>	F <sub>1,28</sub>
Leaf area (cm <sup>2</sup> )								
Betn	0.98 ± 0.02	1.08 ± 0.02	0.98 ± 0.02	1.04 ± 0.03	1.07 ± 0.03	<b>7.96*</b>	1.38	0.55
Salp	3.20 ± 0.14	4.26 ± 0.24	3.85 ± 0.23	3.38 ± 0.19	4.01 ± 0.35	<b>22.7***</b>	<b>5.42*</b>	<b>6.68*</b>
Ledp	0.27 ± 0.01	0.43 ± 0.01	0.35 ± 0.02	0.33 ± 0.02	0.37 ± 0.02	<b>146***</b>	1.02	<b>6.73*</b>
Vacv	0.39 ± 0.01	0.63 ± 0.02	0.48 ± 0.03	0.51 ± 0.03	0.54 ± 0.02	<b>62.0***</b>	0.85	0.51
Specific leaf area (cm <sup>2</sup> · g <sup>-1</sup> )								
Betn	133.3 ± 2.1	158.5 ± 3.3	144.7 ± 3.7	148.1 ± 4.2	144.9 ± 3.2	<b>19.3***</b>	0.24	0.22
Salp	122.5 ± 2.5	125.5 ± 3.5	122.2 ± 3.8	122.6 ± 3.9	127.3 ± 3.7	0.19	0.00	0.84
Ledp	54.6 ± 1.1	62.1 ± 1.3	56.2 ± 1.5	56.8 ± 1.7	62.1 ± 1.2	<b>10.2**</b>	0.04	3.42
Vacv	59.5 ± 1.5	80.4 ± 1.9	68.4 ± 2.5	69.0 ± 2.3	72.6 ± 2.6	<b>71.7***</b>	0.00	1.40
Leaf dry matter content (g · g <sup>-1</sup> )								
Betn	0.55 ± 0.01	0.45 ± 0.01	0.51 ± 0.02	0.49 ± 0.02	0.50 ± 0.02	<b>14.8***</b>	0.67	0.25
Salp	0.50 ± 0.01	0.44 ± 0.02	0.48 ± 0.02	0.47 ± 0.02	0.45 ± 0.02	2.00	0.01	0.36
Ledp	0.54 ± 0.01	0.48 ± 0.01	0.52 ± 0.01	0.51 ± 0.01	0.49 ± 0.01	<b>38.6***</b>	0.60	2.30
Vacv	0.53 ± 0.01	0.47 ± 0.01	0.51 ± 0.01	0.51 ± 0.01	0.49 ± 0.01	<b>12.7**</b>	0.10	1.60
Leaf nitrogen content (%)								
Betn	24.2 ± 0.5	32.8 ± 0.8	28.3 ± 1.0	29.1 ± 1.2	28.0 ± 1.2	<b>61.6***</b>	0.33	0.63
Salp	16.6 ± 0.6	22.4 ± 0.7	20.6 ± 0.7	18.3 ± 1.0	19.9 ± 1.1	<b>26.2***</b>	2.58	1.63
Ledp	14.4 ± 0.5	18.2 ± 0.4	15.5 ± 0.6	16.4 ± 0.6	16.9 ± 0.8	<b>27.3***</b>	1.09	0.22
Vacv	7.8 ± 0.2	11.0 ± 0.6	8.6 ± 0.4	9.3 ± 0.6	10.2 ± 0.8	<b>28.0***</b>	1.03	1.47
Leaf phosphorus content (mg <sup>1</sup> · g <sup>-1</sup> )								
Betn	2.05 ± 0.08	3.90 ± 0.19	2.95 ± 0.24	2.94 ± 0.24	3.03 ± 0.27	<b>60.6***</b>	0.00	0.09
Salp	1.57 ± 0.12	1.55 ± 0.01	1.52 ± 0.10	1.32 ± 0.09	1.81 ± 0.17	0.00	0.17	3.92
Ledp	1.02 ± 0.05	1.32 ± 0.05	1.07 ± 0.06	1.15 ± 0.05	1.29 ± 0.09	<b>13.9**</b>	0.63	2.02
Vacv	0.59 ± 0.03	0.80 ± 0.04	0.64 ± 0.04	0.68 ± 0.05	0.75 ± 0.05	<b>21.8***</b>	0.39	1.63
Leaf carbon to nitrogen ratio								
Betn	20.9 ± 0.4	15.8 ± 0.6	18.2 ± 0.7	18.1 ± 0.9	18.8 ± 0.9	<b>28.5***</b>	0.02	0.33
Salp	29.0 ± 1.4	22.6 ± 0.8	24.0 ± 0.9	27.7 ± 1.8	25.3 ± 1.8	<b>10.9**</b>	2.26	1.32
Ledp	38.0 ± 0.9	29.0 ± 0.7	35.3 ± 1.4	33.3 ± 1.2	32.1 ± 1.2	<b>53.0***</b>	1.91	0.57
Vacv	66.6 ± 1.9	49.7 ± 2.0	62.0 ± 2.7	58.8 ± 3.0	54.1 ± 2.7	<b>45.6***</b>	1.03	2.38

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**Figure 1: Study species: *Betula nana* (a), *Salix pulchra* (b), *Ledum palustre* (c) and *Vaccinium vitis-idaea* (d).**

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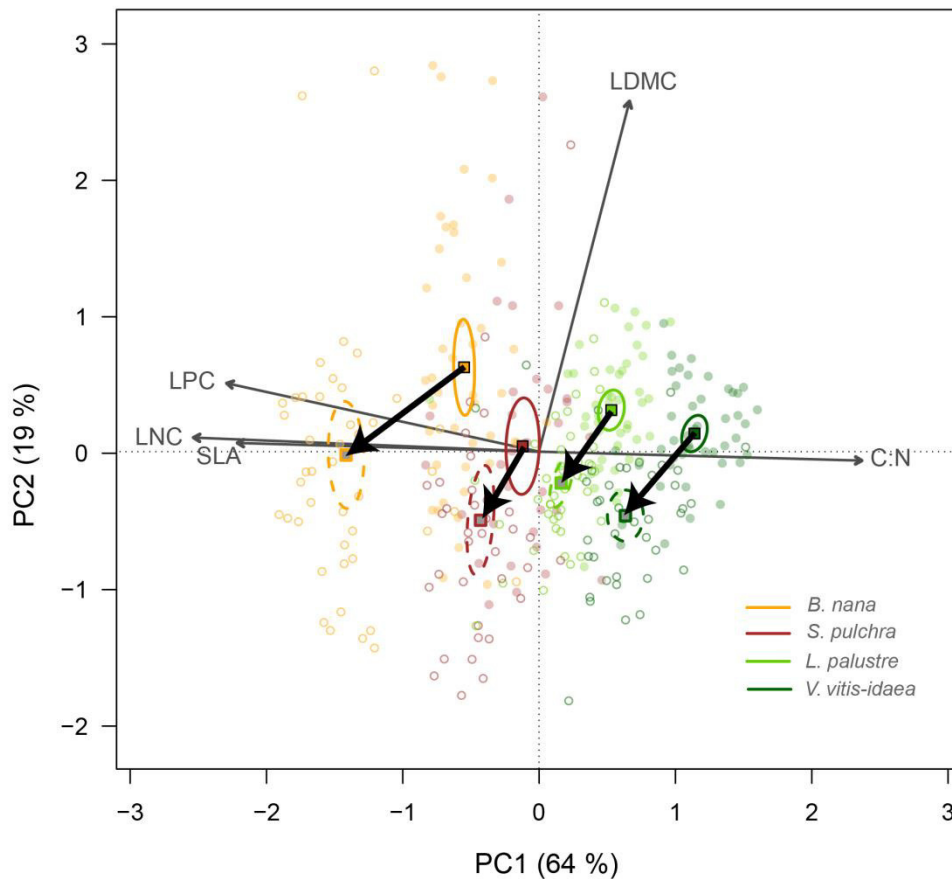
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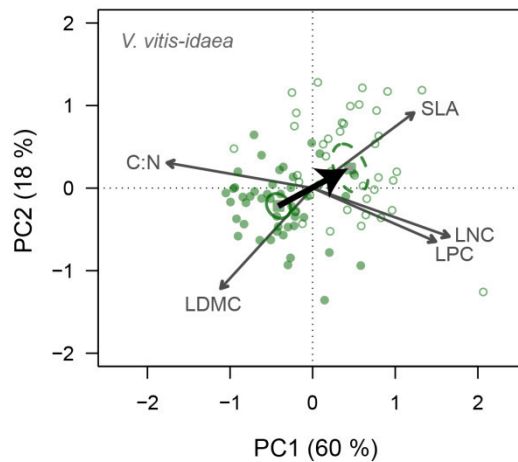
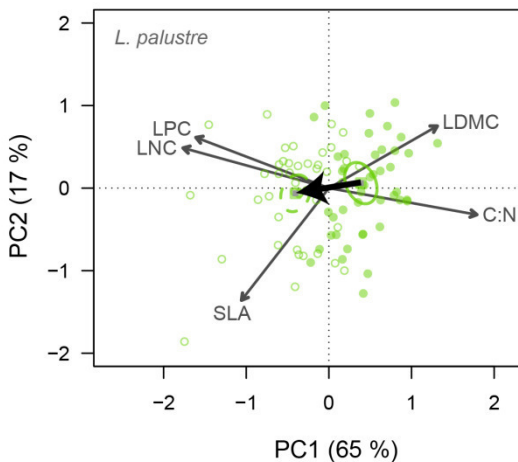
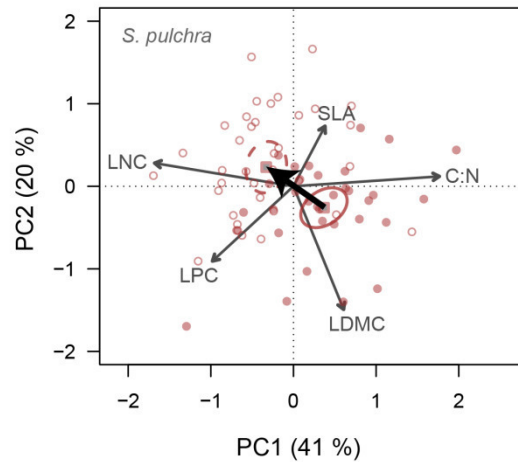
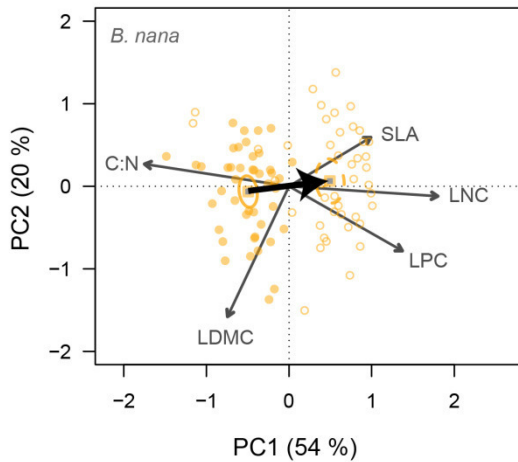
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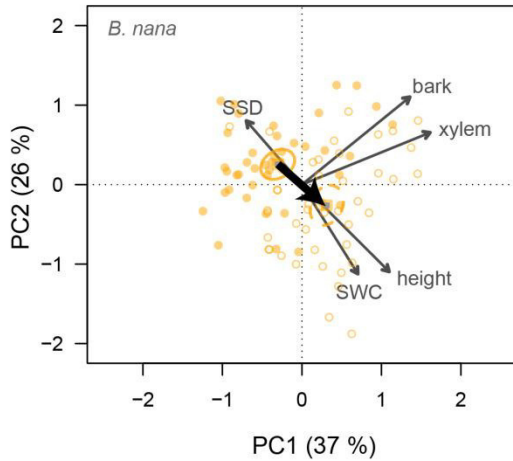
**Figure 2: Principal component biplot of leaf traits for all four shrub species combined, showing change along the leaf conservative-acquisitive continuum (black thick arrows) when nutrients were added. Leaf traits included are leaf dry matter content (LDMC), carbon to nitrogen ratio (C:N), leaf nitrogen content (LNC), leaf phosphorus content (LPC) and specific leaf area (SLA). Points are the trait scores of individuals without fertilization (closed circles) and with fertilization (open circles). Sample scores are scaled by factor 15 and variable loadings by factor 7. Squares indicate the centre of the ordiellipses (standard error with 95% confidence interval) of trait scores without nutrient addition (solid lines) and with nutrient addition (dashed lines). The first principal component explains 64% of the total variance, while the second component explains 19%.**

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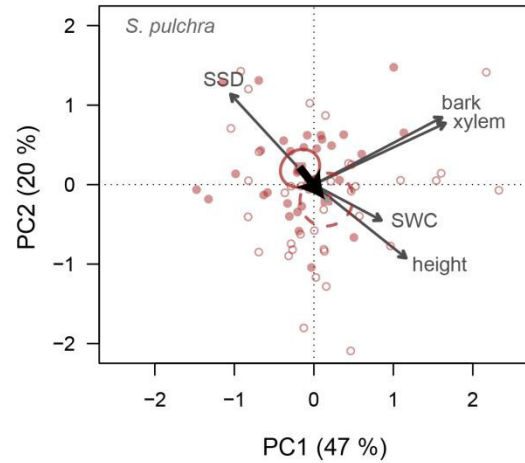


**Figure 3: Principal component biplots of leaf traits for each shrub species. Change in leaf traits when nutrients are added is shown by black thick arrows. Leaf traits included in the PCA are leaf dry matter content (LDMC), carbon to nitrogen ratio (C:N), leaf nitrogen content (LNC), leaf phosphorus content (LPC) and specific leaf area (SLA). Points are the trait scores of individuals without fertilization (closed circles) and with fertilization (open circles). Sample scores are scaled by factor 15 and variable loadings by factor 7. Squares indicate the centre of the ordiellipses (standard error with 95% confidence interval) of the trait scores without nutrient addition (solid lines) and with nutrient addition (dashed lines). The total variance explained by the two first principal components (PC1, PC2) is indicated as percentage between brackets on the axes.**

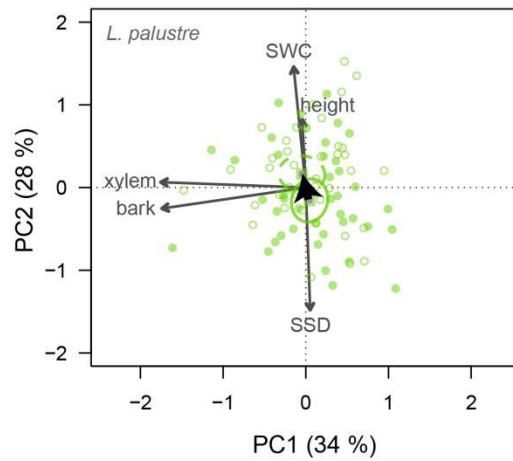
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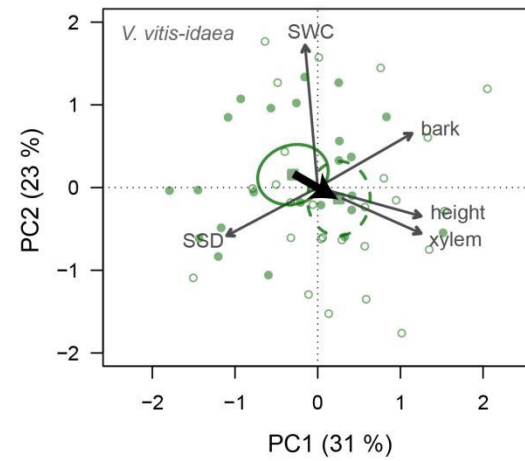
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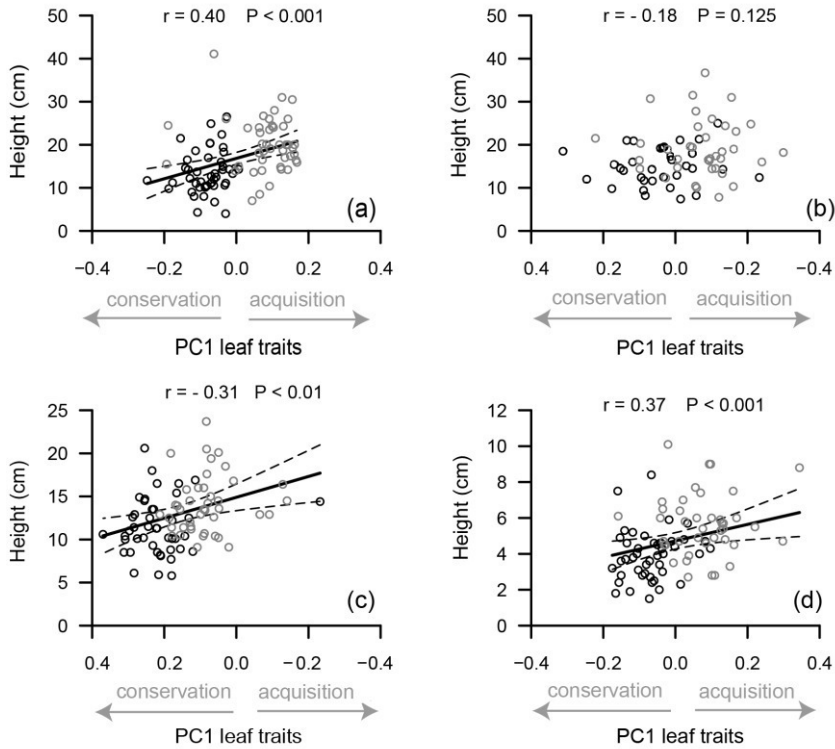
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**Figure 4: Principal component biplots of plant height and stem traits for each shrub species. Stem traits included are xylem diameter, bark thickness, stem water content (SWC), and stem-specific density (SSD). Change in traits when nutrients are added is shown by black thick arrows. Points are the trait scores of individuals without fertilization (closed circles) and with fertilization (open circles). Sample scores are scaled by factor 15 and variable loadings by factor 7. Squares indicate the centre of the ordiellipses (standard error with 95% confidence interval) of the trait scores without nutrient addition (solid lines) and with nutrient addition (dashed lines). The total variance explained by the two first principal components (PC1, PC2) is indicated as percentage between brackets on the axes.**

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**Figure 5: Relationship between plant height and the first axis of the principal component analysis (PC1) of leaf traits for *Betula nana* (a), *Salix pulchra* (b), *Ledum palustre* (c) and *Vaccinium vitis-idaea* (d). Points are trait values for individuals on unfertilized (black) and fertilized plots (grey). Solid lines are values predicted by the linear mixed-effect model and dashed lines are the upper and lower limits of the predicted value confidence interval. Pearson's correlation coefficient ( $r$ ) and p-value ( $P$ ) are indicated on each panel. Main leaf traits comprising PC1 are indicated by grey arrows on the x-axis and grouped into leaf resource acquisition (higher SLA, LNC and LPC) and conservation traits (higher LDMC and leaf C:N).**

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