One leaf for all: Chemical traits of single leaves measured at the leaf surface using near-infrared reflectance spectroscopy

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Abstract

1. The leaf is an essential unit for measures of plant ecological traits. Yet, measures of plant chemical traits are often achieved by merging several leaves, masking potential foliar variation within and among plant individuals. This is also the case with cost-effective measures derived using near-infrared reflectance spectroscopy (NIRS). The calibration models developed for converting NIRS spectral information to chemical traits are typically based on spectra from merged and milled leaves. In this study, we ask whether such calibration models can be applied to spectra derived from whole leaves, providing measures of chemical traits of single leaves.

2. We sampled cohorts of single leaves from different biogeographic regions, growth forms, species and phenological stages to include variation in leaf and chemical traits. For each cohort, we first sampled NIRS spectra from each whole, single leaf, including leaf sizes down to Ø 4 mm (the minimum area of our NIRS application). Next, we merged, milled and tableted the leaves and sampled spectra from the cohort as a tablet. We applied arctic–alpine calibration models to all spectra and derived chemical traits. Finally, we evaluated the performance of the models in predicting chemical traits of whole, single leaves by comparing the traits derived at the level of leaves to that of the tablets.

3. We found that the arctic–alpine calibration models can successfully be applied to single, whole leaves for measures of nitrogen ($R^2 = 0.88$, RMSE = 0.824), phosphorus ($R^2 = 0.65$, RMSE = 0.081) and carbon ($R^2 = 0.78$, RMSE = 2.199) content. For silicon content, we found the method acceptable when applied to silicon-rich growth forms ($R^2 = 0.67$, RMSE = 0.677). We found a considerable variation in chemical trait values among leaves within the cohorts.

4. This time- and cost-efficient NIRS application provides non-destructive measures of a set of chemical traits in single, whole leaves, including leaves of small sizes. The application can facilitate research into the scales of variability of chemical traits and include intra-individual variation. Potential trade-offs among chemical
traits and other traits within the leaf unit can be identified and be related to ecological processes. In sum, this NIRS application can facilitate further ecological understanding of the role of leaf chemical traits.

KEYWORDS

arctic-alpine, carbon, functional groups, leaf chemical traits, nitrogen, phenological stages, phosphorus, silicon

1 | INTRODUCTION

The essential role of chemical constituents in plants and ecosystem functioning is repeatedly emphasized (Aerts & Chapin, 2000; Cebrian et al., 2009; Elser et al., 2007; Elser, Dobberfuhl, MacKay, & Schampel, 1996; Elser, Fagan, Kerkhoff, Swenson, & Enquist, 2010; Fay et al., 2015; Güsewell, 2004; LeBauer & Treseder, 2008; White, 1993). Foliar chemical constituents show interspecific variability at both spatial and temporal scales within and across ecosystems, and are closely related to plant performance and ecological interactions (Güsewell, 2004). Furthermore, foliar chemical content is among the plant traits with the highest intraspecific variability (Albert et al., 2010; Fajardo & Siefert, 2016; Siefert et al., 2015) that may also include intra-individual variability (Ely, Burnett, Lieberman-Cribbin, Serbin, & Rogers, 2019). For instance, trait measures at the leaf level have been found to explain a considerable part of trait variability among tropical trees (Messier, McGill, & Lechowicz, 2010), large leaved food plants (Ely et al., 2019) and alpine plants (Albert et al., 2010). Albert et al. (2010) found intra-individual trait variation to be the largest for leaf dry matter content (LDMC; ratio of dry to fresh leaf mass) in the dwarf shrub Vaccinium myrtillus. LDMC variability of 8%, 37% and 55% was explained by differences among populations, individuals and leaves respectively. However, most methods to measure foliar chemical content require more than a single leaf, especially when working with small arctic and alpine plant species, causing knowledge about intraspecific variability in chemical traits to be at a rudimentary stage. Yet, the single leaf is a plant unit involved in ecological interactions. Thus, chemical trait information at the leaf level is likely to prove useful both to eco-physiological understanding of trade-offs within plants and to how chemical traits, their variability and stoichiometry affect ecosystem process rates (Albert, Grassein, Schurr, Vieilledent, &Violle, 2011; Bolnick et al., 2003, 2011). In this study, we address time- and cost-efficient methodology for measuring several chemical traits in single leaves, a scale of potential relevance to many ecological questions.

Near-infrared reflectance spectroscopy (NIRS) has recently been found to provide cost-efficient and accurate measures of leaf chemical traits independent of species, phenotype, ecological context and region (Couture et al., 2016; Murguzur et al., 2019; Petisco et al., 2005; Serbin, Singh, McNeil, Kingdon, & Townsend, 2014; Serbin et al., 2019; Smis et al., 2014). Such cost-efficient measures open avenues for incorporating leaf chemical traits in large-scale ecological studies. This is strengthened by the fact that a single measure, one spectrum, of a sample is enough for predicting several chemical traits. Furthermore, the sampling of a spectrum is non-destructive, causing analysed plant material to be available for further studies of, for example, the content of other constituents or of ecological processes such as decomposition rates. However, typical NIRS applications are based on calibration models of NIRS-derived spectra from dried and milled leaves versus their analytically derived chemical content. To have enough milled leaf material, it is often necessary to merge several leaves, especially from species with small leaves, masking the potential chemical trait variability among leaves. And although cost-efficient, the process time for milling is still a constraint (Couture et al., 2016). Here we ask whether arctic-alpine calibration models for NIRS-based measures of chemical traits (Murguzur et al., 2019; Smis et al., 2014), models that are based on NIRS spectra from dried, milled and tableted leaves, can be applied to NIRS spectra of single, dried, whole leaves and still provide accurate measures of chemical content.

The precision of NIRS calibration models for measures of chemical constituents is dependent on the precision and bias of the analytical techniques from which the chemical constituents are retrieved and the NIRS spectra are fitted (Chodak, 2008). NIRS calibration models can thus only be as precise as the chemical analysis methods upon which they are based (Coates, 2002). Any analytical technique imprecision can reduce the fit between the actual constituent values and the NIRS spectra, whereas still be within the acceptable range that apply to standard method performance for analytical methods (AOAC International, 2016), Because precision requirements are lower for small contents (Horwitz & Albert, 2006), the fit can be poorer for nutrients with small content. Furthermore, any bias, that is, a systematic shift in measured quantity above or below the true content, will reduce the fit with spectra derived from NIRS. It is therefore a great challenge to assess the actual accuracy of NIRS-based measures. Still, NIRS calibration models, such as the arctic-alpine NIRS calibration models, are not inferior to chemical analysis in terms of accuracy and perform well for a range of chemical constituents (Murguzur et al., 2019; Smis et al., 2014). These models are developed for measures of foliar nitrogen, phosphorus, carbon (Murguzur et al., 2019) and silicon content (Smis et al., 2014). We chose to apply these models in this particular study because they provide accurate measures of chemical traits for a range of growth forms and species, at a range of phenological stages in both arctic and alpine environments (Table 1). Hence, they can potentially provide robust measures of chemical content of single leaves from any species in...
TABLE 1 Performance of arctic-alpine calibration models for foliar chemical content (data from Murguzur et al., 2019; Smis et al., 2014). The calibration models are based on samples from nine (five for silicon) functional groups, three levels of phenology, a range of habitats and three biogeographic regions (one region for silicon). Model parameters are presented for external validation of the calibration models along with information about the samples upon which the calibrations were based. Bias is the mean error between predicted values and chemically measured values.

| Chemical trait | No of species (no of samples) | (A) Content (% dry weight) | (B) Model parameters |
|----------------|-------------------------------|--------------------------|----------------------|
|                |                               | Mean Range               | Intercept Slope Bias R² RMSE |
| Nitrogen       | 97 (552)                      | 2.33 0.34–6.01           | 0.09 0.99 −0.08 0.94 0.20 |
| Phosphorus     | 79 (291)                      | 0.21 0.04–0.64           | 0.05 0.77 0.01 0.76 0.05 |
| Carbon         | 96 (424)                      | 46.05 32.56–56.22        | 2.29 0.95 0.14 0.88 1   |
| Silicon        | 29 (442)                      | 0.47 0–9.99              | −0.001 0.95 −0.05 0.90 0.24 |

2 | MATERIALS AND METHODS

2.1 | Leaf sampling

The sampling was conducted on Svalbard, in Finnmark and in Troms (Norway), representing the biogeographic regions of the high-Arctic, the sub-Arctic alpine and the Boreal-alpine, respectively. The sampling in Svalbard was conducted in Adventdalen (78°10′N, 16°05′E), a wide, formerly glaciated valley on the island of Spitsbergen, during the summer of 2016. We sampled leaves in dry heaths, mesic heaths and wetlands, which represent the majority of habitat types found across the archipelago (Elvebakk, 2005). Both Finnmark and Troms belong to the Norwegian part of Fennoscandia. The sampling in Finnmark was conducted in the low alpine zone at 300–400 m a.s.l. at Ifjordfjellet (70°27′N, 27°08′E), during the summer season of 2015. The region is mainly characterized by dwarf shrub heaths (Walker et al., 2005), whereas we sampled leaves mainly from tundra grasslands that typically dominate river plains and that host a wide variety of growth forms. The sampling in Troms was conducted in the low alpine zone at 400–500 m a.s.l. in the mountainous areas surrounding the city of Tromsø (69°40′N, 18°55′E) during the summer of 2017. Additional sampling of senescent leaves and litter, hereafter denoted as leaf litter, was conducted in the fall in 2017 in the boreal forest of Troms at approx. 50–100 m a.s.l.

We collected a total of 1,677 fresh leaves for a total of 97 leaf-cohorts (set of single leaves merged into tablets), and we collected leaf litter for a total of 20 litter-cohorts (without separating between single leaves; Table 2). Within each biogeographic region, the cohorts were collected from different vegetation types, growth forms, species and from different phenological stages. In addition to assessing whether the arctic-alpine calibration models can be applied to spectra derived from whole leaves, we also assessed the number of spectra needed for predicting accurate chemical content in whole, single leaves as a guide to future sampling. For this purpose, we sampled fresh leaves from the Varanger Peninsula in Finnmark (70°N, 30°E) during the summer season of 2018. We sampled 22 single leaves of different leaf sizes from a total of 18 species, representing forbs, grasses and shrubs (Table S1), and sorted them in size classes of small leaves (Ø < 1 cm), medium leaves (Ø between 1 and 3 cm) and large leaves (Ø > 3 cm).

2.2 | Sample processing

All leaves were sampled individually and immediately put in teabags, pressed dry between filter papers for at least 72 hr and then dried at 60°C for at least 24 hr. In a few cases when we did not have immediate access to plant press and oven facilities, sampled leaves were stored as dry as possible, pressed at the latest during the evening of the sampling and finally dried in an oven within 5 days.

Per cohort we sampled leaves for a total of approx. 100 mg, which is a leaf mass large enough for making a tablet. The final number of leaves per leaf-cohort was on average 17.29, but varied dependent on both the leaf size of the species and the biogeographic region (Table 3). First, we sampled NIRS spectra from whole leaves. From
the leaf-cohorts, spectra were sampled separately from each single leaf, whereas from the litter-cohorts, leaves were stacked and NIRS spectra were sampled from the leaves collectively. After sampling of spectra from whole leaves, all leaves within a cohort were merged and milled into fine powder using a ball mill (Mixer Mill, MM301; Retsch GmbH & Co. Haan) and pressed into tablets (Ø 16 mm, 1 mm thick) using a hydraulic press with 4 tonnes of pressure. Finally, we sampled spectra from each tablet.

Because water shows strong absorption patterns in the near-infrared region (Givens, De Boever, & Deaville, 1997), both the whole leaves and the tablets were oven-dried for 2 hr at 60°C to remove any potential water films, after which they were stored in a desiccator at room temperature (approx. 20°C) until the sampling of spectra.

2.3 | Spectral measurements

All spectra were recorded with a portable NIRS spectrometer (FieldSpec 3, Asd Inc.). Spectra of whole leaves were recorded using a custom-made adaptor that can be attached to the ASD Contact probe (Asd Inc.) and allows for measurements of an area as small as Ø 4 mm (Figure 1). The adaptor was made using Delrin, a non-absorptive material similar to that of the original plant adaptor (advice communicated by Asd Inc.). Spectra of tablets were recorded using a similar setup but with an adaptor for an area of Ø 16 mm, exactly matching the size of the tablets (Smis et al., 2014).

Spectra were recorded with monochromatic radiation in the wavelength range of 350–2,500 nm with NIR, SWIR1 and SWIR2 sensors. The spectra were interpolated to 1 nm intervals based on recordings every 1.4 nm in the 350–1,050 nm region and every 2 nm from 1,050 to 2,500 nm. The arctic–alpine calibration models are based on a subset of the wavelength range, and do not include wavelength regions for which the different sensors overlap (i.e. 350–380, 760–840, 1,700–1,800 and 2,450–2,500 nm) and the visible part of the spectrum (380–720 nm). The same subset should be applied when using the arctic–alpine calibration models for prediction, and hence we prepared the spectra accordingly.

The number of sampled spectra per single, whole leaf was dependent on the leaf area, causing a range in spectra sampled (Table 3). Overall, we sampled from 1 to 12 spectra with an average number of 3.53 replicate spectra per leaf and an average number of 61.03 replicate spectra per leaf-cohort, and 14.1 replicate spectra per litter-cohort. For the tablets, the average was 3 and 4 replicate spectra for leaves and leaf litter, respectively. For the assessment
of the number of spectra needed for predicting accurate content of chemical constituents in single leaves, we sampled 10 replicate spectra from small-sized leaves, an average of 14.5 replicate spectra per medium-sized leaves and an average of 32.5 replicate spectra per large-sized leaves. Each spectrum was recorded as absorbance (log 1/R, where R is reflectance) and the chemical content predicted using the arctic–alpine models for nitrogen, phosphorus, carbon (Murguzur et al., 2019) and silicon content (Smis et al., 2014).

2.4 | Assessment of method performance

We used the predictions of the chemical content of the tablets (for which the arctic–alpine calibration models are developed) as blueprint to which the predicted chemical content of whole leaves was compared. For the whole leaves, we first calculated the average predicted chemical content per single leaf, after which we calculated the average content per leaf-cohort. For the leaf litter, we calculated the average predicted chemical content per litter-cohort directly. We compared the chemical content per cohort as predicted from whole leaves to the chemical content of the cohort as predicted from its tablet. We used linear regression models to assess prediction fit (intercept and slope) and prediction accuracy (root-mean-square error of prediction [RMSEP] and coefficient of determination [R2]). All cohorts were included in the linear regression models for the predictions of nitrogen, phosphorus, carbon (Murguzur et al., 2019) and silicon content (Smis et al., 2014). For the assessment of chemical content variation among single leaves within leaf-cohorts, we first corrected predicted values using correction factors achieved from the regression analyses described above. For the chemical constituents where the fit between predicted content from the whole leaves versus the tablets was not 1:1, we applied the intercept and slope as correction factors to adjust the predicted content per leaf. After correction, the predicted phosphorus and silicon content was negative for a few leaves. These leaves were given a minimum value of content equal to 0.01% phosphorus and 0.1% silicon, similar to the lowest values included in the arctic–alpine calibration models (Table 1).

We assessed intra-cohort variation using a subset of the samples. For intra-cohort variation in chemical content we used leaf-cohorts of Bistorta vivipara, the only species represented with cohorts from all the three biogeographic regions as well as several phenological stages per region. For intra-cohort variation in stoichiometric ratios, we used graminoid cohorts sampled from Svalbard in the late season, representing a range of genera for which the predicted chemical content was based on at least four sampled spectra and for which we could include silicon content. We assessed whether stoichiometric ratios would be more accurately predicted using calibration models based on stoichiometric ratios directly. We made a calibration model for the ratio between nitrogen and carbon (Figure S1a). A comparison between the stoichiometric ratios derived from the arctic–alpine calibration models and the new stoichiometric calibration model indicated they were equally precise (Figure S1b), and we proceeded with the arctic–alpine calibration models.

To estimate the number of NIRS spectra necessary to accurately predict the chemical content of single, whole leaves we sampled a minimum of 10 and a maximum of 42 spectra per leaf. First, we predicted the chemical content from each single spectrum using the arctic–alpine calibration models. Next, we averaged these predictions only if this model was equal or better in accuracy to that of the leaf model, otherwise two separate models (i.e. one for leaf-cohorts and one for litter-cohorts) are presented.
between an increasing number of replicates (average of predictions from the two first replicate spectra, the three first replicate spectra and so on up to the maximum number of replicate spectra for each leaf). Finally, we compared these averages by calculating their differences. We repeated this procedure 10 times, from each of 10 randomizations of the order in which the spectra were taken. We plotted the differences in predictions as a function of the number of replicate spectra. Based on a graphical presentation of the differences, we assessed at what number of replicate spectra the difference in predictions levelled off, with differences approaching zero considered the number of spectra required for accurate predictions of chemical content in single, whole leaves.

All statistical analyses were run in the R environment version 3.4.4 (http://www.r-project.org) using ggplot2 for all graphical presentations.

3 | RESULTS

The range of chemical content derived from sampled NIRS spectra of milled and tableted leaves with the Ø 16 mm plant adaptor (Figure 1) was considerable (Table 4A), providing a range in chemical contents for which to pursue the comparison between predictions from whole leaves and tablets.

We found the arctic-alpine calibration models performed well in predicting content of chemical traits of whole leaves. Predictions of chemical content of a cohort when based on spectra sampled from leaves (using a Ø 4 mm plant adaptor) correlated well with that of predictions based on sampled spectra from the same leaves as milled and tableted (the standardized way of preparing leaf material for measurement of chemical content using NIRS; Figure 2; Table 4B). For nitrogen and silicon, we found both leaf- and litter-cohorts were fitting in a common model (Figure 2). For phosphorus and carbon, we found the slope of the litter-cohorts was steeper than that of the leaf-cohorts (Figure 2). Overall, the predicted content from whole leaves differed in range to that of the predicted content from tablets (Figure 2), and the intercept and slope of the regressions deviated from an ideal relationship of 1:1 for all the chemical traits (Table 4). Hence, to achieve actual chemical content predictions from the Ø 4 mm sampled spectra of whole leaves, the initial predictions from the arctic-alpine calibration models must be corrected.

The predicted chemical content of single leaves within Bistorta leaf-cohorts showed a considerable variation (Figure 3). The range in chemical content among leaves within a cohort was particularly large for the cohorts from Svalbard (Figure 3; Figure S2), and with a larger range in chemical content in early as opposed to late season. In general, the range in predicted chemical content among leaves within cohorts was equal to or larger than the range in predicted

| Chemical trait        | (A) Content (% dry weight) | (B) Model parameters |
|-----------------------|---------------------------|----------------------|
|                       | Mean | Range         | Intercept | Slope | $R^2$ | RMSE |
| Nitrogen              | 2.100 | 0.032–4.515  | 1.073     | 0.604  | 0.88  | 0.824 |
| Phosphorus            |      |               |           |       |       |      |
| Leaf-cohorts          | 0.184 | 0.040–0.443  | −0.014    | 0.842  | 0.65  | 0.081 |
| Litter-cohorts        | 0.127 | 0.040–0.291  | −0.003    | 1.165  | 0.56  | 0.053 |
| Carbon                |      |               |           |       |       |      |
| Leaf-cohorts          | 45.35 | 40.07–51.84  | 7.165     | 0.811  | 0.78  | 2.199 |
| Litter-cohorts        | 46.48 | 43.07–55.19  | −14.403   | 1.341  | 0.91  | 1.654 |
| Silicon\(^a\)         | 0.991 | 0–2.489      | 0.612     | 0.699  | 0.67  | 0.677 |

\(^a\)The Silicon model only includes graminoids and horsetails as these are growth forms with higher silicon content.

| Chemical trait        | Table 4 | Murguzur et al., 2019; Smis et al., 2014 and applied in this study (A). Model parameters for the regression analysis between chemical content predicted from single leaves and tablets (B). For phosphorus and carbon, the model parameters improved when separating leaf- and litter-cohorts, whereas for nitrogen and silicon the best model included both cohorts.

**FIGURE 2** Relationships between nitrogen, phosphorus, carbon and silicon content of whole leaves and tablets, separate for leaf-cohorts (leaf) and litter-cohorts (litter) when their separation improved the linear regression models. The correlation for silicon content is based on silicon-rich growth forms only. The grey line shows the ideal 1:1 relationship
content among seasons and biogeographic regions, as indicated by the predicted content of tablets. The single, whole leaf predictions were attained on the basis of several spectra sampled per leaf but for a few leaves from the Bistorta leaf-cohorts of the Finnmark and Troms regions that were based on one spectrum only (Figure 3). The predicted content of tablets was both larger, similar and smaller than that of the average of predicted chemical content of the single leaves (Figure 3; for all cohorts see Figure 2). In particular, there was a large discrepancy between leaves and tablets for the predicted content of Carbon in the Bistorta leaf-cohorts from the Troms region, the region with most leaves with only one replicate spectrum.

The predicted stoichiometric ratios of single leaves within the graminoid cohorts from Svalbard showed a considerable variation (Figure 4). In general, the three cohorts of grasses showed most variation among leaves, and especially the grass Calamagrostis, where
the variation in the ratio among chemical constituents ranged from seven-fold to 40-fold. The sedge *Eriophorum* showed the least variation among leaves but for the silicon/nitrogen ratio, where it showed the largest variation. For most cohorts, the average of predicted stoichiometric ratios of the leaves overlapped or were close to that of the tablets.

The precision in predicting chemical content per leaf was dependent on the number of sampled spectra. There was a sharp increase

**Figure 5** Differences in predicted foliar nitrogen, phosphorus, carbon and silicon content (% dry weight) as a function of the number of sampled NIRS spectra per leaf, displayed separately for large-, medium- or small-sized leaves. Each dot represents the average difference in prediction, obtained from randomizing the order of the spectra sampled from a leaf and calculating the differences 10 times (with differences between one and two spectra presented as two sampled spectra, differences between two and three spectra as three sampled spectra and so on). The density curves demonstrate the overall pattern across all leaves within a leaf-size group. For a comparison to the scale of the chemical content of the leaves, insets show the average chemical content per leaf-size group. Examples of species representing the different leaf-size groups are the grass *Phleum alpinum* and the forbs *Trollius europaeus* and *Solidago virgaurea* for large-sized leaves, the grasses *Anthoxanthum nipponicum* and *Calamagrostis phragmitoides* and the forbs *Bistorta vivipara* and *Rumex acetosa* for medium-sized leaves, and finally the dwarf shrubs *Vaccinium myrtillus* and *Betula nana* for small-sized leaves. In total, the leaves of 18 species were included (Table S1)
in precision already at 4–5 sampled spectra per leaf, as indicated by a sharp decrease in difference in predictions between 2–3 and 4–5 sampled spectra (Figure 5). When comparing the difference in predicted chemical content to that of the average chemical content of the leaves (insets), the maximum prediction inaccuracy was up to 12.5% when using only two replicate spectra and dropped to approx. 3% when using five spectra. This supports that a few spectra only provide an accurate prediction of foliar chemical content of single leaves.

4 | DISCUSSION

This study shows that the foliar content of a range of key chemical elements can be measured, using NIRS, from a whole, single leaf and from leaf sizes as small as Ø 4 mm. The NIRS application is both time- and cost-efficient, and is non-destructive. In a time when marked changes to the environment are happening, and especially in arctic and alpine regions where predicted changes to biogeochemistry are considerable (Jonasson, Chaplin, & Shaver, 2001), we believe our efficient method to achieve chemical traits is a welcomed contribution (Halbritter et al., 2019).

This is further supported in terms of reduced sampling impact on vegetation when for instance working in experimental or long-term monitoring plots. Also, the quick measure of a range of foliar chemical traits at the level of single leaves opens avenues for research. For instance, chemical traits can be related to that of other traits along with their trade-offs at the level of single leaves, and compared to trade-offs at the level of individuals and populations. Inter- and intra-individual variability in foliar chemical traits can become levels of investigation when studying ecosystem processes such as herbivory and decomposition. In sum, ecological questions for which chemical traits in single leaves are relevant can easily be addressed through our NIRS application.

The arctic-alpine calibration models are based on spectral data of milled plant material pressed into tablets. The purpose of milling and tableting leaf material is to create a homogeneous surface and reduce random light scattering (Smis et al., 2014). Reduced precision is found when predicting from fresh leaves as opposed to dried and milled leaves, yet the loss in precision does not make predictions from fresh leaves inferior (Couture et al., 2016). Furthermore, any gain in information acquired from having time to process more samples (when avoiding tedious processing of leaf samples) may compensate for less accurate predictions (Couture et al., 2016). In our study, any reduced accuracy in the prediction of chemical trait values from the dried and pressed leaves compared to that of the tablets could not be estimated directly as most single leaves were too small for tableting (and too small for providing wet chemistry measures of the chemical content). However, we found that the accuracy of the measure of chemical content in a single leaf increased with the number of spectra sampled, suggesting the uneven surface of a leaf (such as that of veins and other structures) does not interfere with predictions as long as several spectra are sampled.

The arctic-alpine calibration model of silicon performs best for silicon-rich growth forms (Smis et al., 2014). Perhaps for this reason we found the model was only applicable to single leaves of silicon-rich growth forms. Furthermore, species of growth forms with low silicon content such as forbs, shrubs and trees made up two-thirds of all samples (Table 2), hence their inclusion would have caused a bias in the regression analysis towards small content. Also, the predicted silicon content of single leaves of these silicon-poor species was sometimes spurious. This indicates aspects of the leaf surface, in turn affecting the spectral signature, interfered with the Silicon model and that foliar silicon content of silicon-poor growth forms are best measured in a homogeneous surface such as that of milled material (Smis et al., 2014).

The ability to address chemical traits of single leaves provides the opportunity to assess intraspecific chemical trait variability at several scales, including the within individual variability (Albert et al., 2011; Bolnick et al., 2011). Indeed, the nitrogen content related trade-off, or dilemma, of leaves being palatable and efficient in production as opposed to investing in defences (Díaz et al., 2016), may play out differently among single leaves within a plant individual. For instance, plant-herbivore interactions between trees and large ungulates can promote changes at both the modular and genetic levels (Danell, Bergström, Edenius, & Ericsson, 2003). In response to herbivory by moose, the deciduous tree Betula pendula allocates more nitrogen to leaves on shoots browsed by the herbivore than to leaves on lesser-browsed shoots (Danell, Huss-Danell, & Bergstrom, 1985). There are also several other ways by which intraspecific trait variation—that is variation both within and among individuals of the same species—could alter community structure or dynamics (Bolnick et al., 2011). For instance, ecological interactions may depend nonlinearly on the variations in a trait, or trait variation may determine the number of ecological interactions taking place (Bolnick et al., 2011), hence knowledge of the intra-individual trait variation may increase our predictive ability of ecological interactions. In turn, variation in chemical or stoichiometric traits among leaves is likely to drive differences in biodiversity among individual plants. The arctic-alpine calibration models, and potentially also other NIRS-based calibration models, provide an opportunity to address such within-individual variation for a range of chemical traits. Furthermore, NIRS-based spectral information at the leaf level also hold the potential for being scaled up to larger scales. Measures at the leaf level within individual plots can be scaled up to canopy, community and landscape levels, and even larger scales, where for each level confounding factors that blur understanding can be addressed. Such scaling also provide efficient measures of biodiversity (Cavender-Bares et al., 2017).

There are several aspects of leaves for which a focus on their chemical content may be worthwhile. Leaves are functional units for photosynthesis. Leaves are modular units constantly produced and
discarded from plant individuals. Leaves are the units often selected for by herbivores. All these functional roles of leaves suggest their chemical content varies, and that measuring their chemical traits at the scale of the functional leaf unit opens avenues to what questions we can ask in ecology. The arctic-alpine calibration models for NIRS-based prediction of foliar nitrogen, phosphorus, carbon (Murguzur et al., 2019) and silicon content (Smis et al., 2014) can be applied to achieve chemical traits from single, whole leaves, and as such may be the method to open these avenues. Merging the arctic-alpine calibration models with existing calibration models from other regions may further open these avenues towards a global level.

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CONFLICT OF INTEREST

There are no competing financial interests associated to this study.

AUTHORS’ CONTRIBUTIONS

M.P.B., K.A.B., T.M. and S.K. conceived and developed the study; M.P.B. drafted the study design and study protocol; H.B. and K.A.B. did the modelling; K.A.B. drafted the paper with help of M.P.B. and H.B. All authors gave final approval for publication.

DATA AVAILABILITY STATEMENT

Data are accessible at UiT Open Research Data (https://opendata.uit.no, https://doi.org/10.18710/MHX1Z7).

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Additional supporting information may be found online in the Supporting Information section.

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