Assessing the efficiencies and challenges for nutrient uptake by aquatic plants

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Abstract
Aquatic plant meadows are valuable components to the ‘coastal filter’ and it is important to understand the processes that drive their ability to cycle nutrients. However, at present, the field-based evidence for understanding the drivers of nutrient uptake by plants is lacking. This study aimed to investigate how well individual shoots of aquatic plants could meet their nitrogen demands using the sediment nutrient pool (porewater ammonium) and to explore which traits helped to facilitate such uptake. Several species were investigated in shallow, submerged (2-4 m) mixed-species communities in the northern Baltic Sea using incubation experiments with enriched ammonium. After a 3.5 h incubation time, individuals were collected and analysed for nitrogen (% DW) and $^{15}$N (at-%) concentrations. Uptake by plants was calculated per unit nitrogen in response to the $^{15}$N-labelled source and to overall nitrogen availability. Background porewater ammonium availability was highly variable between individual plants. Species identity did not significantly affect uptake metrics and the effect of ambient porewater availability was weak. As biomass increased there were significant logarithmic declines in the 95th quantiles of nutrient uptake rates, ambient porewater nutrient availability and aboveground nitrogen tissue concentrations (% DW). Such findings suggested that uptake rates of plants were significantly demand-driven and the nutrient conditions of the porewater were significantly driven by the demands of the plant. Findings parameterised the unfulfilled potential for some aquatic plants to cycle nutrients more efficiently and highlighted the potential importance of access to new nutrient sources as a way of enhancing nutrient cycling by aquatic plants. Plant traits and community properties such as the activity of infauna could facilitate such an access and are likely important for nutrient uptake.
Introduction

Aquatic plant meadows are valuable for cycling nutrients in coastal zones (McGlathery et al., 2007). They absorb dissolved nutrients and store them as biomass, which is then available for further processes such as burial or grazing by consumers, and it ultimately slows the re-release of inorganic nutrients into the surrounding environment (McGlathery et al., 2007). Understanding uptake processes is particularly important in the Baltic Sea because many of its ecosystems are threatened by eutrophication (see Andersen et al., 2009; Gustafsson et al., 2012) which has been accelerated by an increase in nutrient concentrations from anthropogenic nutrient loading to coastal zones (Gustafsson et al., 2012). It is hence imperative to investigate how well aquatic plants can access and utilise nutrient pools in the surrounding environment and understand what enhances their ability to cycle nutrients.

Aquatic plants absorb dissolved organic and inorganic nutrients from the water column and, mainly, the sediment porewater (Erftemeijer and Middelburg, 1995; Touchette and Burkholder, 2000). Nutrient availability in the sediment is increased by processes such as excretion by fauna (Peterson and Heck, 2001; Peterson and Heck, 1999) as well as the decomposition of organic matter (Kenworthy and Thayer, 1984), including vegetation and fauna, phytodetritus (Evrard et al., 2005) and other particles entrained from the water column (Kennedy et al., 2010). Such factors are highly spatiotemporally variable (e.g. Hemminga, 1998; Wirachwong and Holmer, 2010; Fourquarean et al., 1992). Nutrients can be depleted to the extent that aquatic plant growth is nutrient-limited (Perez et al., 1991; Ferdie and Fourquarean, 2004; Armitage et al., 2011); which is not always the case (as observed by Olsen...
and Valiela, 2010; Mutchler and Hoffman, 2017; Campbell and Fourqurean, 2014), but nutrient availability can drive spatial differences in plant patch development (Furman et al., 2017). Such a large-scale response to nutrient availability is owing to the responsive foraging and growth strategies by roots and clonal ramets to nutrient availability (Furman et al., 2017; Kembel et al., 2008; Campbell et al., 1991; de Kroon and Mommer, 2006). Thus, nutrient variability is highly influential to the development of aquatic plant meadows and their ability to cycle nutrients.

If an individual plant had a greater capacity to meet its nutrient demands using the resources that were available (e.g. Comas and Eissenstat, 2004), it would cycle nutrients more efficiently in the place that it grows, be more robust to temporal change in nutrient availability and in turn the meadow structure would be more robust to changes in nutrient availability. The growth of the root network of a plant is coordinated by its hormonal cues in response to nutrient demands. In terrestrial environments, the growth responses and root architectural strategies to meet nutrient demands change between species (Kembel et al., 2008; Campbell et al., 1991). These strategies can enhance uptake to different levels of success depending on the environment. For instance, having a higher root length per unit root mass (Specific Root Length, SRL) reduces the distance that nutrients must diffuse to reach the roots, and this benefits plants in nutrient poor environments because it catalyses nutrient supply (Aerts, 1999). However, in nutrient-rich zones the nutrient supply is no longer the limiting factor; instead it is the rate at which plants can absorb the available nutrients and so their physiological nutrient uptake kinetics become more important for enhancing nutrient uptake (Aerts, 1999). In the context of aquatic plant communities, we do not know which strategies are valuable for uptake and thus, what the nutrient microenvironment is like for an individual plant. While the physiology of different aquatic plant species can vary (e.g. Gustafsson and Norkko, 2016), coarser-scale morphological variations which affect uptake
rates are still yet to be explored. For instance, seagrasses and other aquatic plants often develop a network of ramets which can share nutrients (Marbà et al., 2002); a strategy which enhances nutrient uptake in terrestrial environments with heterogeneous nutrient availability (Roiloa and Hutchings, 2013). Yet for any single shoot in natural conditions, we do not understand which conditions are most important for uptake success. For seagrass, root traits are likely to represent extremely poor nutrient conditions because the spatiotemporal variability of nutrients can be so influential on meadow growth (Furman et al., 2017) that the variability of traits which enhance uptake kinetics are likely to be redundant compared to the ability for plants to increase their reach to access new nutrient sources. In this case, growing longer roots would likely be the most important attribute for nutrient uptake rather than the amount of absorptive surfaces (i.e. Root biomass). However, without evidence we do not know whether nutrient uptake by each seagrass shoot is dictated by the abundance of nutrients available to be absorbed, whether plants benefit from investing more in extracting nutrients from their immediate environment (e.g. High SRL) or the ability for roots to access new nutrient pools (e.g. Maximum root length) is the most important trait in the natural environment.

As nitrogen is hypothesised to be the main nutrient which limits growth at higher latitudes (Hemminga, 1998; Short, 1987), and sediment porewater ammonium the main nitrogen source, porewater ammonium was the focal nutrient of this study. Our aim was to quantify nitrogen uptake rates by several aquatic plant species from the sediment porewater environment. We hypothesised that species would have significantly different nutrient uptake rates, and these differences would be owing to the variation in morphological root traits between species. It would be highly difficult to conduct such an investigation in typical seagrass meadows because they are often monocultures (Hemminga and Duarte, 2000), which means that there would not likely be enough variability in uptake strategies to find
insightful trends. In the northern Baltic Sea, the brackish-water conditions allow for a unique assembly of marine, estuarine and limnic species to coincide within the same meadow (Kautsky, 1988; Gustafsson and Norkko, 2016), therefore these communities were ideal for capturing a variety of uptake strategies facilitated by interspecific differences, while environmental conditions in the experiment remained relatively consistent. In this experiment, we examined how efficiently individual shoots could fulfil their nitrogen demand in response to a nutrient enrichment, and by comparing uptake rates between several species, we explored which uptake strategies were most beneficial to uptake rates. It is important to investigate which strategies enhance uptake in aquatic plants because it allows us to identify which mechanisms might otherwise be limiting nutrient uptake, thus nutrient cycling, in aquatic plant communities in the northern Baltic Sea.

Methods

Study site and vegetation

The experiment was conducted in the shallow, submerged (2-4 m) mixed species vegetative communities around Tvärminne Zoological Station, Finland. Tvärminne is situated on the Hanko Peninsula of the Finnish archipelago, Baltic Sea (59° 50' 40" N, 23° 14' 56" E WGS84). The species measured both have a limnic and marine origin; the freshwater species *Ceratophyllum demersum*, *Myriophyllum spicatum*, *Stuckenia pectinata*, *Potamogeton perfoliatus* and *Zannichellia major*, and the marine/brackish species *Zostera marina* and *Ruppia cirrhosa*. We incubated independent ramets of a clone to prevent translocation during incubation. Individuals with a single shoot were selected, however for small species with a trailing growth (*R. cirrhosa* and *Z. major*), incubating multiple shoots was unavoidable. To prevent multiple species from being in the same incubation, we focussed on sparse stands. While sparse stands may not facilitate optimal conditions for aquatic plants, there is also less
competition for nutrients and it is possible to investigate the nutrient uptake drivers of a
single plant shoot without the influence of other shoots around. All incubations were at least
1 m apart from each other and individuals did not have noticeable grazer marks nor
biofouling. Species-averaged canopy heights can range from 10-70 cm in this area and season
(Gustafsson and Norkko, 2016), but individuals in this study did not exceed 20 cm in height.
The sediment of the study site had approximately 0.5-0.8 % OM content and 1.2-7.3
% silt fraction (<0.063 mm) (Gustafsson and Norkko, 2018).

Field work

Short term nutrient enrichment incubations were conducted *in situ* using SCUBA (Figure 1).
$^{15}$N - enriched ammonium sulfate solution (40 ml, 47 μM, 99 at-%) was used as an
immediate-release sediment fertiliser followed by incubations (Figure 1). 36 incubations were
conducted on 4 days, 9 incubations on each day, during the late growth season (August-
September) 2015 and 10 further incubations were conducted in September 2016 (Table 2). A
minimum of 3 replicates of each species were incubated on each day and a further 3
specimens were collected intact from the surrounding meadow as ambient samples to obtain
the background $^{15}$N in plant tissue. Not all incubations were successful due to insufficient
root or belowground biomass available for the analysis, but this remained unknown until
harvest (see Table 2). Rooting depths can vary between species (approx. 5-20 cm, pers. obs.)
and within species (length can change by a magnitude of 10, Gustafsson and Norkko, 2018),
which meant that selecting a standardised depth for enriching the sediment porewater and for
measuring the ambient porewater nutrient availability to plants required due consideration.
For instance, injecting the liquid fertiliser too shallow into the sediment would likely lead to
an immediate re-release of the majority of fertiliser into the water column. Likewise, too deep
an enrichment and the fertiliser would be out of reach of roots for some of the individuals.
The sediment porewater was enriched at 7-8 cm deep; approximately half way down the depth of the core and close to the maximum lengths that the roots of aquatic plants in the study area typically reached (Gustafsson and Norkko, unpubl.). At this depth, it was predicted that the nutrients would reach the plants by diffusion and they would not be immediately diffused into the water column (Cheng et al., 2014, Chowdhury and Bakri, 2006; Clavero et al., 2000). The plants absorbed a fraction of the enriched source, meaning that there was enriched solution available for the plants throughout the experiment (Figure 1). Porewater samplers collected depth-integrated water samples in the upper 10 cm of sediment (see Gustafsson and Boström, 2011; Figure 1) to provide a best estimate of nutrient availability to plants in the region where roots were mostly found (pers. obs.) before the enrichment occurred. This data was used to estimate background porewater nutrient availability within the area which varied widely (Table 1).

Water column samples were periodically collected (13/08/15; 10/09/15; 11/09/15; 02/09/16) to estimate ammonium, nitrate, nitrite and phosphate availability to the aboveground parts of the aquatic plants (Table 1).

**Uptake metric calculations**

The increase in nitrogen availability induced by the experimental enrichment was calculated as the percentage of nitrogen added compared to ambient NH$_4^+$ concentrations, in moles, for each individual. Uptake by plants was calculated per unit nitrogen in order to detect the small-scale changes in nitrogen attributed to the short incubation time. The ability for plants to respond to the specific nutrient enrichment (response rates, μgN gN$^{-1}$ h$^{-1}$, hereafter RR) was determined using the equation:

\[ {^{15}\text{N}} \text{(enriched plant)} - {^{15}\text{N}} \text{(ambient plant)} \]
Where the $^{15}\text{N}$ of an ambient plant represented an average of 3 replicates of each species. Uptake rates which considered additional nutrient uptake by the ambient porewater nutrient pool (uptake rates, $\mu\text{gN g}^{-1} \text{ h}^{-1}$, hereafter UR) were calculated using mass-balanced isotope mixing equations (Collos, 1987) as described by Lepoint et al. (2002). The important difference between uptake response rate (RR) and uptake rate (UR) is that RR is a measurement of uptake of the enriched source, but UR incorporates ambient porewater ammonium measurements to estimate total ammonium uptake during the incubation. To avoid confusion, both units of uptake rate are named uptake metrics when they are being referred to collectively.

Statistical analysis

Analyses were performed using the statistical software R (version 3.02.2, R Core Team 2015). For all analyses, the relevant assumptions relating to heteroscedasticity and the distribution of residuals were checked and in order to meet the criteria, some of the variables (RR, UR, porewater NH$_4^+$, total biomass, root biomass) were logarithmically transformed.

The effect of total biomass on uptake metrics, NH$_4^+$ and leaf tissue nitrogen concentrations were analysed using 95$^{\text{th}}$ quantile regression. Quantile regression, such as 95$^{\text{th}}$ quantile analysis, is an effective technique to explore the maximum (or factor ceiling) effect that a factor has on a dependent variable (Thomson et al., 1996; Thrush et al., 2003). 95$^{\text{th}}$ quantiles were calculated for 50mg biomass bins, which were then analysed using linear regression. Thus, statistical outputs represent trends of the boundary lines for the 95$^{\text{th}}$ quantiles. One individual was removed from biomass-specific analyses because of its inordinately high biomass compared to the rest of the samples. The 2016 data was pooled with 2015 data for quantile regression analyses, but not for analyses which required species
averages because only two species were incubated in 2016 and their pooled values may have
led to misrepresentative differences between species.

A multiple-regression style General Linear Model (GLM) was used to analyse the
effect of species identity, ambient porewater ammonium availability (log-transformed,
hereafter NH$_4^+$) and their main effects on uptake metrics. For species comparisons, 3
replicates of each species were randomly selected from 2015 data for equal group sizes,
however Ceratophyllum demersum was discounted because it did not have true roots and
another species (Zannichellia major) was removed due to replicate loss.

Morphological characteristics of the experimental plants were not measured because it
was important to process plant samples as quickly as possible after harvesting in order to
prevent $^{15}$N-loss from samples through leaching. Instead, the 95$^{th}$ quantile regressions were
used to infer which traits were likely to be important to enhance uptake rates, and pre-existing
data was compared to uptake rates as a supportive tool for highlighting opportunities for
future study. Species-averaged root traits were derived from a plant survey in the same study
area at the same time of year, the previous year (Gustafsson and Norkko, 2018). These traits
represented relative species differences in root traits for individuals growing in the study area
(Table 3); from up to 5 quadrats per species, wherein up to 10 of the longest roots were
measured. While rooting depth can vary between individuals, by using maximum root length
the data represented the length that the roots of each species could typically extend to within
the study area. Then, individual effects of NH$_4^+$ and root biomass on uptake metrics were
analysed using linear regression. Species-averaged root traits (Mean Max Root Length,
MMRL; Specific Root Length, SRL) were tested against uptake metrics with rank regression
using up to three randomly selected uptake rates for each species. Z. major, which had been
previously discounted from the species comparison, was included in the trait analyses
because it did not risk violation of the test assumptions and its presence did not change the significant outcome of the tests.

Results

Porewater nutrient availability

Background porewater ammonium availability was highly variable within the replicates of individual plants (Table 1), which meant that when plants were experimentally enriched with identical doses of fertiliser the effect on porewater nutrient availability varied widely (approximately 2-74 % NH$_4^+$ enrichment, based on 20% sediment porosity measured at the site) and the starting nutrient conditions were not consistent between individuals. Such porewater variability had a significant, yet weak effect on uptake metrics (Table 4). Uptake response rates (RR) and uptake rates (UR) exhibited large variation too; ranging from -1.31 and -30.15 μgN gN$^{-1}$ h$^{-1}$ respectively, where uptake of the enriched source did not exceed the natural variability of $^{15}$N, to 109.43 and 1032.25 μgN gN$^{-1}$ h$^{-1}$. It was highly unlikely that the nutrient enrichment produced toxic effects for the plants because the enrichment was minor in comparison to the variability of ambient nutrients in the sediment porewater.

Biomass and nutrient cycling

There were no trends between root biomass and uptake metrics (Table 4), however total biomass was significantly linked to the upper limits of all the nutrient cycling metrics that were measured in this study. For instance, there were significant logarithmic declines in the 95$^{th}$ percentiles of uptake metrics (UR & RR) as biomass increased (Figure 2A & B, Table 5), and larger individuals were more likely to be limited in their overall uptake rate (UR) rather than their capacity to respond to the experimental enrichment (RR, Figure 2A & B, Table 5). Similarly, the sediment porewater surrounding larger plants was significantly less likely to be NH$_4^+$ replete (Figure 2C, Table 5) and leaf tissue nitrogen concentrations of larger plants did
not reach the same maximum concentrations as for smaller plants; a trend which was not
singly controlled by species identity (Figure 2D).

Root traits and uptake rates

Species did not have significantly different uptake rates (UR) nor response rates (RR) ($t_{17, 14} = .654, p > .05$; $t_{17, 14} = .765, p > .05$, respectively), nor did species identity interact with porewater ammonium availability to produce significantly different URs and RRs ($t_{17, 14} = -.998, p > .05$; $t_{17, 14} = -.881, p > .05$). Supportive analyses which refer to an external trait dataset suggest that while there may not have been species differences, species traits could provide insights into the drivers of uptake rates. For instance, species capable of growing longer roots (Mean Max Root Length, MMRL) were significantly more adept at responding to the experimental nutrient pulse by rank (RR; Figure 3A, Table 4) but MMRL was not significantly beneficial for the rank order of overall uptake rates (UR; Figure 3B, Table 4). There were no significant relationships between Specific Root Lengths (SRL) and the ranks of uptake metrics (UR & RR, Table 4).
This study has provided novel insights into the short-term dynamics of nutrient uptake by aquatic plants, not only for plants in the northern Baltic Sea but also for vascular plants in other submerged aquatic ecosystems. Overall, results showed that the nutrient demands of the plants significantly affected uptake rates, nutrient storage in plant tissue and the nutrient availability in the sediment porewater (Figure 2, Figure 4). Therefore, short-term nutrient uptake in these aquatic plant meadows was likely to be largely driven by the presence and characteristics of the aquatic plants, more so than by environmental nutrient availability. For example, as plants increased in size, their ability to cycle nutrients to their full potential was significantly hindered and the nutrient availability in the sediment porewater was likewise depleted (Figure 2). This suggests that access to new nutrient sources is likely to benefit nutrient uptake and nutrient cycling by plants. Such conclusions are likewise suggested by supportive analyses which identify a potential link between maximum root lengths and uptake rates (Table 4). These insights are informative and beneficial because they described the challenges for nutrient uptake experienced by a single aquatic plant shoot, and findings parameterise the unfulfilled potential for some aquatic plants to cycle nutrients more efficiently.

Unlike other characteristics of a plant, such as root length and thickness, biomass does not only represent the capacity for plants to acquire nitrogen (e.g. Tessier and Raynal, 2003) but it also represents their nitrogen demand (Poorter and Nagel, 2000). As an individual increases in size, the amount of nitrogen it needs to maintain the same nutrient concentrations in its more abundant tissues becomes exponentially higher (Figure 2, Blackman, 1919). While the overall nitrogen uptake rates of the plants in this study were likely to be higher as plants increased in biomass (hereafter ‘larger plants’) their capacity to meet their increased demand...
was significantly limited (Figure 2A & B). Thus, the increased capacity to absorb nitrogen that a larger biomass might provide, such as by having more absorptive surfaces (Blackman, 1919), was outweighed by the increased demand from having larger standing nitrogen stocks (Figure 2). Also, larger plants were less likely to store the same standing nitrogen stocks (%DW) as smaller plants (Figure 2D), suggesting that larger plants may not be absorbing nitrogen to their full potential. This is a new insight into the efficiency of nutrient cycling by aquatic plants. In addition, the porewater nitrogen was more likely to be depleted in the vicinity of larger plants (Figure 2C). All trends were highly variable for plants with a smaller biomass, which made factor ceiling analysis a highly valuable tool. Its results suggest that firstly, demand was represented more strongly than supply in biomass-uptake relationships (Figure 2A & B), secondly the larger plants did not store N (%DW) standing stocks as high as smaller plants, which was not necessarily related to species (Figure 2D) and lastly an individual plant can deplete porewater NH$_4^+$ (Figure 2C). Due to the aforementioned observations, the evidence suggests that the larger plants were nitrogen deprived. Such nitrogen deprivation may not have limited growth because plants are capable of luxury nutrient uptake (e.g. Göransson and Eldhuset, 1991) and the threshold N (%DW) concentration above which further uptake becomes luxurious is unknown. Nevertheless, larger plants were likely to not have been cycling nitrogen to their fullest potential and even larger plants that may be growing naturally at higher densities than the selection criteria of this study are probably more N-deprived. Thus, nitrogen is likely to be a competitive resource in aquatic plant meadows of the north Baltic Sea, and interpretations of species interactions during nitrogen uptake should be re-evaluated.

Aquatic plants are subject to a nutrient compromise in the water column (Fourquean et al., 1992); that with increasing nutrient availability in the water column, the light availability decreases as responsive phytoplankton blooms attenuate the light which is
transmitted to the plants from the surface. But results from this study exhibited an additional nitrogen compromise for plants that might not be expected in a eutrophicated sea (Gustafsson et al., 2012). Ammonium concentrations in the sediment porewater were lower than the global median for aquatic plants meadows (Hemminga, 1998). Trends showed that overall, the more investment in biomass the less capable plants were at meeting their nutrient demands (Figure 2D), which has been previously unobserved in the field. These trends between aquatic plant biomass and tissue nitrogen concentrations represented not only the ability for plants to absorb their nutrients from the sediment, but also the water column, and the trade-off between investing in belowground and aboveground biomass for uptake is also accounted for. Therefore, these interpretations apply to the capacity for a whole plant to meet its nutrient demands. It is noteworthy that this experiment was conducted at the peak for plant biomass, but not for production, and so if the experiment was conducted at the peak of productivity the biomass-uptake trends observed could become even more accentuated. Overall, these findings show that larger aquatic plants are not cycling nutrients to their full potential. Thus, they could potentially become more efficient at their role in the coastal nutrient filter if circumstances allowed it. Strategies, such as morphological root traits, could thus be highly valuable for helping plants to cycle nutrients more effectively. For instance, depending on the extent of nutrient variability, the ability for plants to absorb nutrients in the vicinity (i.e. Root biomass, Specific Root Length) or to access new nutrient pools (i.e. Maximum root length) could be highly valuable for enhancing nutrient uptake.

Root traits and uptake rates

In general, an increase in ambient nutrient availability leads to an accelerating increase in the rate of nutrient uptake by an individual, typically described by a hyperbolic trend (Caassen and Barber, 1976). However, such a trend was not detected in this experiment. Porewater estimates may not have been representative of the relative availability of nutrients in the
depletion zones, or an interaction with other factors that were more influential drivers of uptake rate (e.g. Plant traits, plant biomass, Figure 4) could have convoluted the conventional porewater nutrient-uptake trend. This was likewise the case for root biomass (Table 4), because root biomass represents a dynamic equilibrium between environmental nutrient availability and plant nutrient demands, making it highly unlikely that a direct relationship could be observed between root biomass and any plant nutrient parameters (e.g. Ruffel et al., 2011; Figure 4). Given that the uptake rates, plant stores and sediment supply of nutrients were largely influenced by the plant biomass in this study, the nutrient dynamics were likely to be largely plant-driven. These findings suggest that access to new nutrient sources would enhance uptake rates and plant stores. Similarly, when uptake rates were compared to pre-existing trait data, species with the potential to grow longer roots appeared to be more equipped to respond to the nutrient source (Figure 2). This seemed less beneficial when the effect of porewater availability was also taken into account to estimate overall uptake rates (Figure 2). Thus, the plants would potentially benefit from longer roots to respond to the nutrient enrichment, but because longer roots cannot guarantee a proportional increase in access to more nutrients it does not necessarily mean that the overall uptake rate would be faster. Within the zone from which plants can deplete nutrients, it is unlikely that properties that increase the absorptive properties of the roots would be as helpful as accessing new zones with nutrient sources. Otherwise, one would expect that porewater nutrient availability or root biomass would have had stronger roles in the different uptake rates of individuals of this study (Figure 2; Figure 4). Instead, the plant biomass (i.e. Demand) had the prevailing influence (Figure 2). Supporting this result, specific root length was not significantly linked to the rank order of uptake rates in the supportive analyses. It is unlikely that variability of further physiological strategies for absorption would significantly affect uptake rates. Therefore, findings imply that even for a single aquatic plant shoot, the variability in access
to a nutrient source is more important than variability for plants to retrieve the nitrogen once it had been accessed. Other traits which might enhance access to nutrient sources include expanding the spatial reach for nutrients than seeking nutrients within a smaller zone, based on terrestrial examples (see Kembel et al., 2008; Campbell et al., 1991 and references within), which could include growing multiple ramets because labour is divided between ramets, simultaneously increasing the reach (Roiloa and Hutchings, 2013).

Conclusion

To our knowledge, this is the first study to measure the nitrogen uptake of a variety of different aquatic species in relatively similar environmental conditions in the field. Furthermore, the influence of different plant characteristics was surveyed in a natural environment while other factors remained relatively similar. Results show a previously unidentified, extreme spatial variability of sediment porewater ammonium availability within an aquatic plant meadow, and its variability is reflected by the uptake strategies, challenges and successes for even a single aquatic plant shoot. Overall, our results suggest that the larger plants could perform better at their role as nutrient filters in the coastal zone, and increasing access to new nutrient sources could help plants to uptake and store nutrients more effectively. Thus, larger plants have the potential to cycle nutrients more effectively, and this could potentially be facilitated by increasing the access to new nutrient pools which would benefit from nutrient cycling by aquatic plants. These findings fill a knowledge gap because such conditions for aquatic plants in their natural settings were previously unknown. Future studies which investigate the mechanisms that increase root access to new nutrient sources could help to understand how nutrient cycling by aquatic plants might be enhanced. For example, more diverse aquatic meadows may enhance resource partitioning (Gustafsson and Boström, 2011) and thus nutrients may be cycled more effectively, and nutrient cycling by fauna can increase plant growth (Peterson and Heck, 2001; Peterson and Heck, 1999). To
follow from this study, there is an opportunity to investigate how community composition
(Gustafsson and Boström, 2011), the activity of infauna (e.g. Villnäs et al., 2018) and other
factors increase the connectivity of aquatic plant nutrient depletion zones to areas which
would benefit from their role as nutrient filters. Ultimately, such studies may help to develop
effective conservation strategies to mitigate eutrophication by providing the most effective
solutions to enhance the role of aquatic plant meadows as nutrient filters.

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Helsinki.

Literature cited


**Tables**

Table 1. Overall mean, standard deviation (SD) and frequency (n) of ammonium (NH$_4^+$), nitrate and nitrite (NO$_2^-$ + NO$_3^-$) and phosphate (PO$_4^{3-}$) concentrations (µg L$^{-1}$) in the sediment porewater next to individual plants and the water column.

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<th></th>
<th>Mean</th>
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<th>N</th>
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<td><strong>Porewater</strong></td>
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<td>NH$_4^+$</td>
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<td>NO$_2^-$ + NO$_3^-$</td>
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<td>PO$_4^{3-}$</td>
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Table 2. Details of enrichment events including dates, which species were enriched (Treatment), number of successful replicates for each species (originally 3 replicates per species per day in 2015. No replicates lost in 2016), average temperature (°C) and photosynthetically active radiation (PAR: mol m$^{-2}$ h$^{-1}$; HOBO Pendant® Temperature/Light Data Logger 64 K, Onset, USA, light data converted to PAR as described by Gustafsson, Norkko 2016) during each incubation.

<table>
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<th>Enrichment</th>
<th>Date</th>
<th>Treatment</th>
<th>Successful replicates</th>
<th>Temp. °C</th>
<th>PAR (mol m$^{-2}$ h$^{-1}$)</th>
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<td>6</td>
<td>02.09.16</td>
<td>Myrspi, Potper</td>
<td>7, 3</td>
<td>16</td>
<td>1.1</td>
</tr>
</tbody>
</table>
Table 3. Mean Max Root Length (mm) and Specific Root Length means (± Standard Error, SE) of study species in the same area of the experiment. Data source Gustafsson & Norkko (2018).

<table>
<thead>
<tr>
<th>Species</th>
<th>MMRL (mm)</th>
<th>SRL</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Myriophyllum spicatum</em></td>
<td>9.8 ± 1.3</td>
<td>5.7 ± 1.6</td>
</tr>
<tr>
<td><em>Potamogeton perfoliatus</em></td>
<td>7.4 ± 0.4</td>
<td>14.0 ± 1.2</td>
</tr>
<tr>
<td><em>Ruppia cirrhosa</em></td>
<td>8.7 ± 1.2</td>
<td>11.6 ± 1.3</td>
</tr>
<tr>
<td><em>Struckenia pectinata</em></td>
<td>6.4 ± 1.2</td>
<td>19.5 ± 2.0</td>
</tr>
<tr>
<td><em>Zannichellia majus</em></td>
<td>7.5 ± 1.4</td>
<td>11.2 ± 0.6</td>
</tr>
<tr>
<td><em>Zostera marina</em></td>
<td>6.4 ± 0.8</td>
<td>8.5 ± 0.7</td>
</tr>
</tbody>
</table>

Table 4. Regression statistics for aquatic plant uptake rate (UR, μgN gN⁻¹ h⁻¹) and uptake response rate (RR, μg¹⁵N gN⁻¹ h⁻¹), in response to changes in porewater ammonium availability (NH₄⁺, μM) and root biomass (mg). Rank regression statistics for UR and RR with changes in species-averaged Mean Max Root Length (MMRL, mm) and species-averaged specific root length (SRL).

<table>
<thead>
<tr>
<th></th>
<th>Regression statistics</th>
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<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Coefficient</td>
<td>Intercept</td>
<td>r²</td>
<td>n</td>
<td>F</td>
</tr>
<tr>
<td><strong>Porewater [NH₄⁺]</strong></td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>UR</td>
<td>.9112</td>
<td>3.5734</td>
<td>.2902</td>
<td>30</td>
<td>11.45</td>
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<tr>
<td>RR</td>
<td>.2556</td>
<td>2.3424</td>
<td>.0694</td>
<td>30</td>
<td>2.088</td>
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<tr>
<td><strong>Root biomass (mg)</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
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<tr>
<td>UR</td>
<td>-.007912</td>
<td>6.185049</td>
<td>.02568</td>
<td>17</td>
<td>.3954</td>
</tr>
<tr>
<td>RR</td>
<td>-.002668</td>
<td>2.204714</td>
<td>.002517</td>
<td>23</td>
<td>.05299</td>
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<table>
<thead>
<tr>
<th></th>
<th>Rank regression statistics</th>
<th></th>
<th></th>
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<tbody>
<tr>
<td></td>
<td>Coefficient</td>
<td>Intercept</td>
<td>n</td>
<td>Overall Wald test</td>
<td>P</td>
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<tr>
<td><strong>Mean max root length</strong></td>
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<tr>
<td>UR</td>
<td>262.63</td>
<td>-1570.02</td>
<td>6</td>
<td>3.5724</td>
<td>.053</td>
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<tr>
<td>RR</td>
<td>4.0781</td>
<td>-214567</td>
<td>6</td>
<td>4.2730</td>
<td>.034</td>
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<tr>
<td><strong>Specific root length</strong></td>
<td></td>
<td></td>
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<tr>
<td>UR</td>
<td>-.69579</td>
<td>182358</td>
<td>6</td>
<td>1.3207</td>
<td>.296</td>
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<tr>
<td>RR</td>
<td>-29.799</td>
<td>657346</td>
<td>6</td>
<td>0.6716</td>
<td>.526</td>
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</tbody>
</table>
Table 5. 95th quantile regression statistics for aquatic plant uptake rate (ln(UR), μgN gN⁻¹ h⁻¹), uptake response rate (ln(RR), μgN gN⁻¹ h⁻¹), aboveground nitrogen concentration (AG N, %DW) and porewater ammonium availability (NH₄⁺, μM) when compared to total biomass (ln(biomass), mg) of an individual plant.

<table>
<thead>
<tr>
<th></th>
<th>Coefficient</th>
<th>Intercept</th>
<th>r²</th>
<th>N</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total biomass</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>UR</td>
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<td>18.0999</td>
<td>0.774</td>
<td>12</td>
<td>38.67</td>
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<tr>
<td>RR</td>
<td>-1.5419</td>
<td>11.1518</td>
<td>0.7212</td>
<td>12</td>
<td>29.46</td>
<td>&lt;.0001</td>
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<tr>
<td>AG N (%DW)</td>
<td>-0.8825</td>
<td>7.3764</td>
<td>0.7969</td>
<td>15</td>
<td>55.92</td>
<td>&lt;.0001</td>
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<tr>
<td>Porewater NH₄⁺</td>
<td>-12.380</td>
<td>93.398</td>
<td>0.5943</td>
<td>12</td>
<td>17.11</td>
<td>.002</td>
</tr>
</tbody>
</table>

Figure legends

Figure 1. Experimental design of an incubation for a single aquatic plant. Multiple incubations (min. batch of 9) were conducted simultaneously within an incubation day.

Figure 2. Total plant biomass (ln(biomass), mg dry weight (DWT)) and, A: Uptake rates (ln (UR), μgN gN⁻¹ h⁻¹), B: Uptake response rates (ln (RR), μgN gN⁻¹ h⁻¹), C: Porewater ammonium (NH₄⁺, μM) and D: Aboveground nitrogen concentration (AG N, percentage dry weight; %DW). ‘D’ displays data for enriched and non-enriched plants. Dotted lines denote 95th percentiles. 95th percentiles were calculated before the data was logarithmically transformed; two values are missing from A and B due to disproportionately negative values.

Figure 3. Species-averaged mean max root lengths (mm) and (A) uptake response rates (ln(RR+10)), (B) species-averaged uptake rates (ln(UR+100)), (μgNgN⁻¹h⁻¹).

Figure 4. Conceptual diagram of the trade-offs occurring in nutrient uptake for aquatic plants in the northern Baltic Sea, based on current evidence from this study and published literature. Solid lines:
Evidence-based trends considering interspecific and intraspecific trends. Dashed line: May not apply to a single shoot of a plant that has multiple ramets (Roiloa, Hutchings 2013). Reverse trend for plants with multiple ramets. *(= Marba et al. 2002, while other components to the model were inferred from the results of this study.)*