

Toxicity of reduced nitrogen in eelgrass (*Zostera marina*) is highly dependent on shoot density and pH

T. van der Heide · A. J. P. Smolders · B. G. A. Rijkens ·
E. H. van Nes · M. M. van Katwijk · J. G. M. Roelofs

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Abstract In sheltered, eutrophicated estuaries, reduced nitrogen (NH_x), and pH levels in the water layer can be greatly enhanced. In laboratory experiments, we studied the interactive effects of NH_x , pH, and shoot density on the physiology and survival of eelgrass (*Zostera marina*). We tested long-term tolerance to NH_x at pH 8 in a 5-week experiment. Short-term tolerance was tested for two shoot densities at both pH 8 and 9 in a 5-day experiment. At pH 8, eelgrass accumulated nitrogen as free amino acids when exposed to high loads of NH_x , but showed no signs of necrosis. Low shoot density treatments became necrotic within days when exposed to NH_x at pH 9. Increased NH_3 intrusion and carbon limitation seemed to be the cause of this, as intracellular NH_x could no longer be assimilated.

Remarkably, experiments with high shoot densities at pH 9 showed hardly any necrosis, as the plants seemed to be able to alleviate the toxic effects of high NH_x loads through joint NH_x uptake. Our results suggest that NH_x toxicity can be important in worldwide observed seagrass mass mortalities. We argue that the mitigating effect of high seagrass biomass on NH_x toxicity is a positive feedback mechanism, potentially leading to alternative stable states in field conditions.

Keywords Alternative stable states · Ammonia · Ammonium toxicity · Density dependence · Necrosis · Positive feedback

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T. van der Heide (✉) · M. M. van Katwijk
Department of Environmental Science,
Institute for Wetland and Water Research,
Radboud University Nijmegen, Faculty of Science,
P.O. Box 9010, 6500 GL Nijmegen, The Netherlands
e-mail: t.vanderheide@science.ru.nl

T. van der Heide · A. J. P. Smolders · B. G. A. Rijkens ·
J. G. M. Roelofs
Department of Environmental Biology,
Institute for Water and Wetland Research,
Radboud University Nijmegen, Faculty of Science,
P.O. Box 9010, 6500 GL Nijmegen, The Netherlands

E. H. van Nes
Aquatic Ecology and Water Quality Management Group,
Department of Environmental Sciences, Wageningen University,
P.O. Box 8080, 6700 DD Wageningen, The Netherlands

Introduction

Seagrass meadows in tropical to temperate coastal areas are among the most productive ecosystems on earth, and harbor high biodiversity. They provide forage and refuge for a vast number marine animal species (Duarte 2002; Orth et al. 2006). At present, these ecosystems are increasingly being lost worldwide, often characterized by mass mortality of seagrasses (Jackson et al. 2001). Multiple stressors have been identified as causes for the decline (e.g., Duarte 2002; Orth et al. 2006). Seagrass losses have been ascribed to climate change (e.g., temperature and sea level rise), sedimentation, disease, turbidity, and toxicity events (e.g., Duarte 2002; Orth et al. 2006). In temperate estuaries, (anthropogenic) eutrophication is one of the most important processes fuelling turbidity and toxicity events (e.g., Goodman et al. 1995; Azzoni et al. 2001; Nielsen et al. 2002b; Kemp et al. 2005; Perez et al. 2007). Seagrass mass mortality in these areas has been linked with high concentrations of sulfide in the sediment pore water near the end of

the growing season often in combination with high turbidity levels in the water layer (e.g., Carlson et al. 1994; Goodman et al. 1995; Azzoni et al. 2001; Pedersen et al. 2004). However, elevated levels of inorganic nitrogen in the water layer may also be important. These compounds have been shown to cause severe toxicity in seagrasses and may therefore help explain these dramatic events (Burkholder et al. 1992; van Katwijk et al. 1997; Bird et al. 1998; Brun et al. 2002).

In this perspective, toxicity caused by reduced nitrogen compounds (NH_x) is particularly interesting, as it is a common phenomenon described for a vast number of aquatic and terrestrial plant species (Britto and Kronzucker 2002). In seagrasses, ammonium (NH_4^+) is taken up directly by both leaves and roots (Iizumi and Hattori 1982; Thursby and Harlin 1982; Rubio et al. 2007). Even though NH_4^+ is preferred over nitrate (NO_3^-) as a nutrient source (Touchette and Burkholder 2000), it has been found to cause severe toxicity within weeks when concentrations become too high (van Katwijk et al. 1997; Bird et al. 1998; Brun et al. 2002). In seagrass beds growing in sheltered eutrophicated estuaries, NH_x concentrations may be greatly enhanced ($>200 \mu\text{mol l}^{-1}$) by degradation processes inside competing macroalgal mats (Hauxwell et al. 2001), discharges of waste and river water (Brun et al. 2002), or due to natural die-off of phytoplankton, macroalgae, or seagrass itself near the end of the growing season (Landers 1982; Farnsworth-Lee and Baker 2000). Additionally, the pH in some of these stagnant lagoons may rise up to 9 or even 10 during daytime due to photosynthesis by the same species (Choo et al. 2002; Feike et al. 2007; our unpublished data). This in turn increases concentrations of highly toxic gaseous ammonia (NH_3), which is converted from ammonium as the pH level rises (e.g., Farnsworth-Lee and Baker 2000; Körner et al. 2001, 2003; Nimptsch and Pflugmacher 2007).

As NH_x is both a valuable nutrient source and a toxic substance for eelgrass, we hypothesize that its toxicity is dependent on the shoot density in meadows. At a certain load, NH_x concentrations in the canopy will most likely be significantly lower in high shoot density meadows compared to beds with low shoot densities, because concentrations will be actively lowered through the direct uptake of NH_x by the leaves (Iizumi and Hattori 1982; Thursby and Harlin 1982). Thus, it can be expected that a certain load may cause severe toxicity in a low density bed whereas the effect will be much less severe or even absent in a high density seagrass bed.

In this study, we investigated the interactive effects of NH_x , pH, and shoot density in the water layer on the physiology and survival of the seagrass *Zostera marina* (commonly called eelgrass). We conducted two laboratory experiments. First, we tested the eelgrass long-term tolerance

for NH_x at natural sea water pH. Secondly, we conducted an experiment to test the interactive effects of NH_x , pH, and shoot density on the short-term survival of eelgrass. Finally, we discuss the relevance of our results with regard to (eelgrass) physiology and ecology.

Materials and methods

After collection in the field, plants were stored at 10°C for transport to the laboratory in Nijmegen (Netherlands). Shoots (aboveground parts attached to at least 4 cm of rhizome) were acclimatized for 2 weeks in 100-l glass containers (l:w:h = 80:30:45 cm) that were placed in a climate controlled room at 20°C . The culture medium used for acclimatization and the experiments was prepared from deionized water and Tropic Marin[®] synthetic sea salt. We used a salinity equal to the level measured in the field. We set light intensity at $180 \mu\text{mol m}^{-2} \text{s}^{-1}$ with a day:night cycle of 16:8 h. This is comparable to average light levels in the field during the growing season around depths of 2–3 m (assuming a light attenuation coefficient of 0.5 m^{-1}) (e.g., van der Heide et al. 2007), which is in the normal depth ranges of eelgrass (e.g., Nielsen et al. 2002a).

At the start of both experiments, we carefully selected healthy apical shoots, based on shoot length, number of leaves, and the percentage mortality (black coloration) of the leaves. Only shoots with four or five leaves and a mortality of less than 5% in the three youngest leaves were used in the experiments. The plants were allowed to acclimatize for two additional days, after placement in the experimental units.

Long-term experiment

The goal of this experiment was to simulate the response of eelgrass to NH_x exposure in ‘open ocean’ conditions. Plants were obtained at the end of August from the Atlantic Ocean near the shores of Brest (France). Based on the literature, we expected the plants to show a strong response to NH_x exposure due to the low ambient nitrogen concentrations ($\pm 11 \mu\text{M}$ total N) and a relatively high salinity in this area (± 33 PSU) (van Katwijk et al. 1999). The experiment was carried out in a 5-week period in 16 glass containers (l:w:h = 20:20:58 cm) placed in a 20°C water bath. Each unit contained 20 l of medium and 8 cm of sandy sediment from the sampling site. Two shoots were placed in each container (total shoot biomass $0.68 \pm 0.04 \text{ g DW}$). The culture medium in the containers was continuously replenished from stock containers at a renewal rate of twice a week, using Masterflex peristaltic pumps. We used four different treatments, resulting in four replicas per treatment. NH_x concentrations in the stock solutions were 20, 60, 100,

and $150 \mu\text{mol l}^{-1}$ by addition of NH_4Cl . We added $1 \mu\text{mol l}^{-1}$ cyanoguanidine to the stock solutions to prevent nitrification of ammonium (Smolders et al. 1996). The pH level was carefully kept between 8.0 and 8.2 by aerating the units with CO_2 enriched air (5 ppt CO_2), using mass flow controllers. The total dissolved inorganic carbon concentration (DIC) in the water was $2.2 \pm 0.02 \text{ mM}$, a close approximation of the DIC level of the sampling site (2.3 mM). Water samples for analysis of NH_x and nitrate were taken on a weekly basis. Leaf growth, leaf number, and shoot biomass were determined before and after the experiment.

Short-term experiment

Alongside the 5-week experiment, we performed a 5-day experiment with eelgrass from the Atlantic Ocean (near Brest, see previous heading), focusing on the acute combined effects of pH and NH_x (setup, pH, NH_x levels, and analyses were performed as described below). In this trial, we found severe necrosis in eelgrass leaves due to NH_x addition at pH 9.1 but not at pH 8.1 (see Fig. S1 in Supplementary material). However, these results pose virtually no ecological relevance as raised pH levels almost only occur in sheltered eutrophicated estuaries with a much lower salinity due to nutrient-rich freshwater influence. Therefore, we repeated this experiment at a lower salinity, using plants collected from the Baltic Sea near Kiel (Germany) at the end of August 2006. We expected this population to respond somewhat more conservatively to NH_x exposure compared to the Brest population, as nitrogen loads in the Baltic Sea are higher ($\pm 27 \mu\text{M}$ ambient total N) and salinity is much lower ($\pm 16 \text{ PSU}$) compared to the waters around Brest (van Katwijk et al. 1999). The experiment was conducted in round glass columns containing 4 l of medium ($\text{Ø:h} = 11:45 \text{ cm}$) placed in a water bath at 20°C . We used four NH_x concentrations (0, 50, 100, and $250 \mu\text{mol l}^{-1}$ added as NH_4Cl) at two pH levels (8.1 and 9.1). Additional to the initial setup of the short-term trial, we used two shoot densities at the highest pH to investigate a possible density dependent effect: 1 or 10 shoots per container, corresponding to 88 and 884 shoots m^{-2} (total shoot biomass 0.44 ± 0.12 and $4.25 \pm 0.28 \text{ g DW}$, respectively). At pH 8, experiments were performed using low shoot densities only, as a much lower toxicity was expected. In total, we used 48 glass columns, resulting in four replicas per treatment. Because of the short experimental period, plants could be maintained without sediment (e.g., Thursby and Harlin 1982; Rubio et al. 2007). This prevented NH_x loss due to nitrification or adsorption to sediment particles. Shoots were positioned in the columns by attaching stainless steel nuts to the rhizomes. As NH_x affinity is much higher for leaves than for roots (Iizumi and Hattori 1982; Thursby and Harlin 1982) and because uptake via roots and

rhizomes is strongly limited in eelgrass when aboveground tissues are exposed to NH_x (Thursby and Harlin 1982), we assumed that NH_x assimilation through the remaining belowground tissue was negligible in our experiment. During the experiment, the culture medium was replenished each day at the start of the light period. The pH was checked regularly during the day and kept between 8.0 and 8.2 or 9.0 and 9.2, by adding HCl or NaOH. DIC of the 16 PSU water was $1.85 \pm 0.03 \text{ mM}$, comparable to the values measured at the sampling site (1.88 mM). Water samples for analysis of NH_x were taken daily just before replenishment.

Analysis of leaf tissue fitness

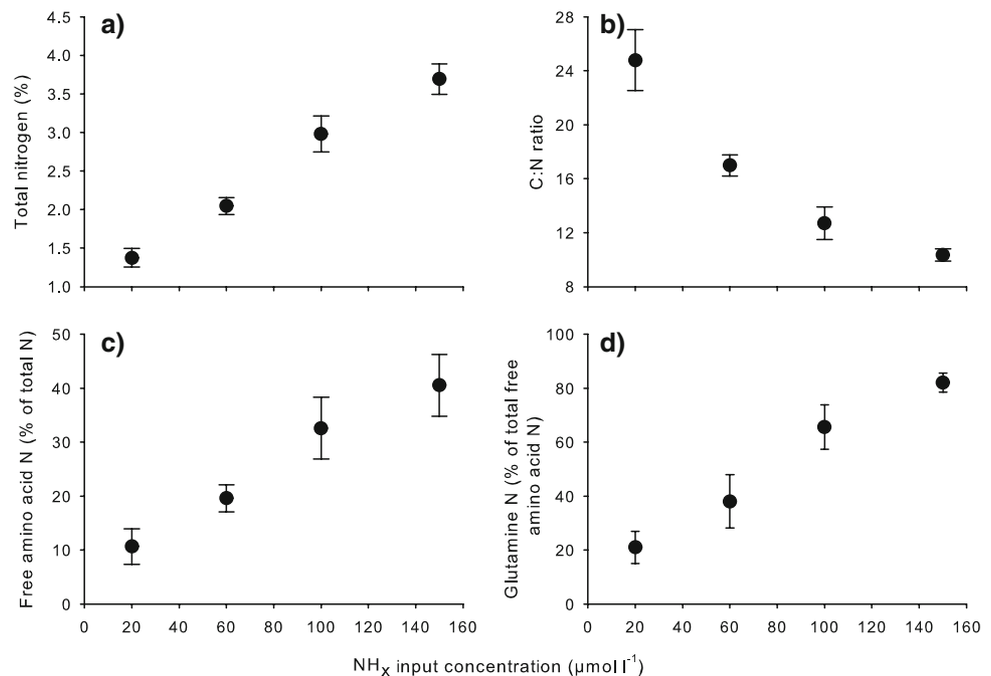
After each experiment, we examined the fitness of the leaves by scanning the three youngest leaves of each shoot at 600 dpi, resulting in a resolution of over half a million pixels per leaf. We subsequently analyzed the scans using Adobe Photoshop. First, we eliminated background noise in the picture using the “threshold” function. The image was then converted into a 64-color index picture (red–green–blue scheme). Next, we used the “color range” function to distinguish between dead and living leaf tissues. The amount of green coloration per pixel was used to determine whether leaf tissue was either dead or alive. Based on tests with fit and dead leaves from the field, we found that pixels with a green value of over 70 (± 2 ; on a scale of 0–255) should be considered alive while lower values could be regarded as dead tissue. Based on this procedure, dead, living, and background areas in the picture were then colored in the three primary screen colors (red, green and blue). Finally, we calculated the percentage of living and dead tissues from the histogram values of the image.

Chemical analyses

Water samples taken during both experiments were analyzed for NH_x and nitrate (Tomassen et al. 2003). NH_x concentrations were determined spectrophotometrically using hypochlorite. Nitrate was measured spectrophotometrically with sulphanilamide after reduction of nitrate to nitrite in a cadmium column.

Free amino acids in the leaves of the shoots were extracted from healthy looking tissues, according to Van Dijk and Roelofs (1988). After extraction, amino acid concentrations were measured after pre-column derivatisation with 9-fluorenylmethyl-chloroformate (FMOC-Cl), using high pressure liquid chromatography (HPLC) (Tomassen et al. 2003). C:N ratios and total N concentrations were determined in freeze-dried leaf tissues by an CNS analyzer (type NA1500; Carlo Erba Instruments, Milan, Italy).

Fig. 1 Effects of four NH_x input concentrations on eelgrass (*Zostera marina*) leaf tissues (mean \pm SE) in the long-term experiment: **a** total nitrogen (N) content, **b** C:N ratio, **c** free amino acid N relative to total N content, and **d** glutamine N relative to the total free amino acid N content



Statistical analyses

For all relevant variables, the effects of NH_x in the long-term experiment and NH_x , pH, density and their interactions in the short-term experiment were tested by analyses of variance (ANOVA) and multifactor ANOVA, respectively. Prior to the analyses, outliers were omitted based on Dixon's Q test (Dean and Dixon 1951) and data were tested for normality. For multiple comparisons of normally distributed means (post hoc tests), we used independent samples t tests. In the long-term experiment, we compared the effect of NH_x between treatments (6 tests). For the short-term experiment, we evaluated the effect of NH_x in all pH and density treatments (18 tests), compared group means of both pH levels at low density (4 tests) and tested the effect of density in the pH nine treatments (4 tests). Thus, 26 hypotheses were tested in total. Type I error in the testing procedures was controlled using the false discovery rate method ($P^* = 0.05$) (Benjamini and Hochberg 1995; Verhoeven et al. 2005).

Results

Long-term experiment

Data were normally distributed and no outliers were detected. Within 1 week, NH_x concentrations in the different treatments stabilized at 1, 12, 38, and 75 (± 2) $\mu\text{mol l}^{-1}$. Nitrate levels were negligible in all treatments during the

entire experimental period. After 5 weeks, we found no significant change in biomass, leaf growth, or in number of leaves. Moreover, shoots in all treatments looked healthy and leaf survival showed no trend at all, being over 90% in all treatments. In contrast, ANOVA revealed significant differences for internal nitrogen concentrations ($F = 34.9$, $P < 0.001$) and C:N ratios ($F = 21.9$, $P < 0.001$) in the leaf tissues. Post hoc test showed that, apart from the two highest loads, all treatments differed significantly from each other for both variables. The total nitrogen content in the leaves ranged from 1.4 to 3.7% from the lowest to the highest treatments, respectively (Fig. 1a). Average C:N ratios varied from 25 in the lowest to 10 in the highest treatments (Fig. 1b).

The amount of nitrogen that was stored as free amino acids in the tissues varied substantially with the applied NH_x load. Concentrations of free amino acid N increased significantly from 113 $\mu\text{mol N/g DW}$ in the lowest NH_x treatment to 1,093 $\mu\text{mol/g DW}$ in the highest treatment (ANOVA, $F = 10.3$, $P = 0.001$). The percentage of total tissue nitrogen stored as free amino acids varied from 11 in the lowest to 41% in the highest treatment (ANOVA, $F = 8.6$, $P = 0.003$; Fig. 1c). Excess nitrogen in the shoots was primarily stored as glutamine. The percentage of glutamine in the total free amino acid concentration changed from 21% in the lowest to 82% in the highest NH_x treatment (ANOVA, $F = 13.9$, $P < 0.001$; Fig. 1d). Post hoc tests revealed significant differences between the lowest and the two highest loads for all three variables. Additionally, the 60 $\mu\text{mol l}^{-1}$ NH_x treatment also differed significantly with the highest load for these variables.

Table 1 Results of the multi-factor ANOVAs on all relevant variables in the short-term experiment

	NH _x treat.	pH	Density	NH _x treat. × pH	NH _x treat. × density
Leaf tissue survival (%)	29.1***	92.1***	82.9***	20.7***	13.6***
Total N (%)	92.8***	4.3*	6.6*	3.3*	2.9 ns
C:N ratio (g:g)	64.8***	0.7 ns	5.9*	0.4 ns	2.2 ns
Free amino acids N of tot. N (%)	313.8***	96.0***	7.9**	46.1***	4.7**
Total free amino acids N (μmol l ⁻¹)	339.6***	112.0***	7.8**	79.2***	5.8**
Glutamine N of tot. amino acids N (%)	339.8***	2.1 ns	49.9***	1.8 ns	18.0***
NH _x conc. after 24 h (μmol l ⁻¹)	3514.4***	11.0**	887.6***	0.622 ns	212.5***

All data were normally distributed, except for total nitrogen. This variable obtained normality after inverse transformation

F values and significance levels are shown for all main effects and their interactions

*0.01 ≤ *P* ≤ 0.05; **0.001 ≤ *P* < 0.01; ****P* < 0.001

Short-term experiment

Analysis showed that all data were normally distributed and that 2 outliers should be excluded. Survival was significantly affected by NH_x, pH, density, and their interactions (ANOVA, Table 1). Leaf tissue survival ranged from 94% in the controls to 57% in the 250 μmol l⁻¹, pH 9 treatment. Necrosis in the leaf tissues was expressed as a typical brown-black discoloration (Fig. 2). Within the pH 8 treatment, there was no significant effect of any of the NH_x loads on leaf tissue survival. At pH 9, NH_x significantly affected survival in both density treatments, but the intensity of the effects was clearly different. In the high density columns, only the highest NH_x treatment deviated significantly from the control treatment, whereas in the low density columns both the 50 and 250 μmol l⁻¹ NH_x treatments differed significantly from the control (and from each other) (Fig. 3a). The effects of pH and density were significant in the 50 and 250 μmol l⁻¹ NH_x treatments.

Total nitrogen content ranged from 1.5% in the control treatments to 2.6% in the 250 μmol l⁻¹, pH 8 treatment (Fig. 3b). The effect of the NH_x treatments was highly significant (Table 1). Multiple comparison tests on the NH_x treatments were significant in 13 out of 16 cases. The effect of pH was only significant in the 250 μmol l⁻¹ treatment, whereas density was significant in the 100 μmol l⁻¹ treatment. The C:N ratio showed a trend that was inversely correlated with the total N content (Fig. 3c). The average ratio ranged from 27 in the control treatments to 15 in the 250 μmol l⁻¹, pH 8 treatment.

The percentage of nitrogen that was stored in free amino acids increased significantly with increased NH_x loads (significant in 14 out of 16 tests). Contents ranged from 3% in the controls to a maximum of 39% in the 250 μmol l⁻¹, pH 8 treatment (Fig. 3d), equaling 36 and 702 μmol free amino acid N/g DW, respectively. The effect of pH was significant in the 50 and 250 μmol l⁻¹ treatments, whereas density showed significant effects in the 50 and 100 μmol l⁻¹

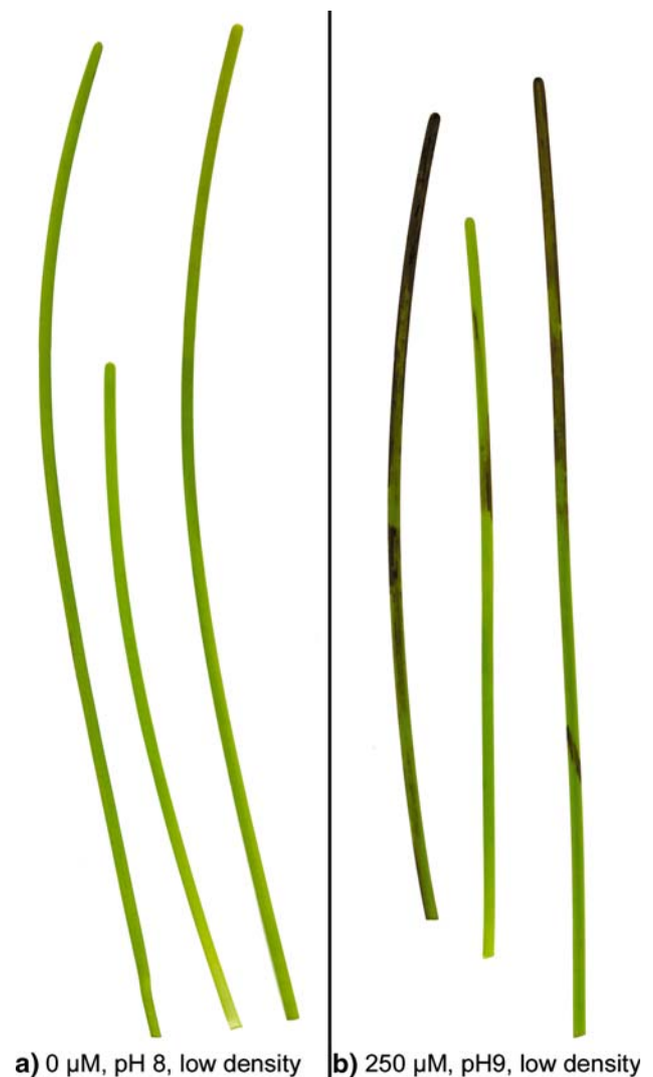
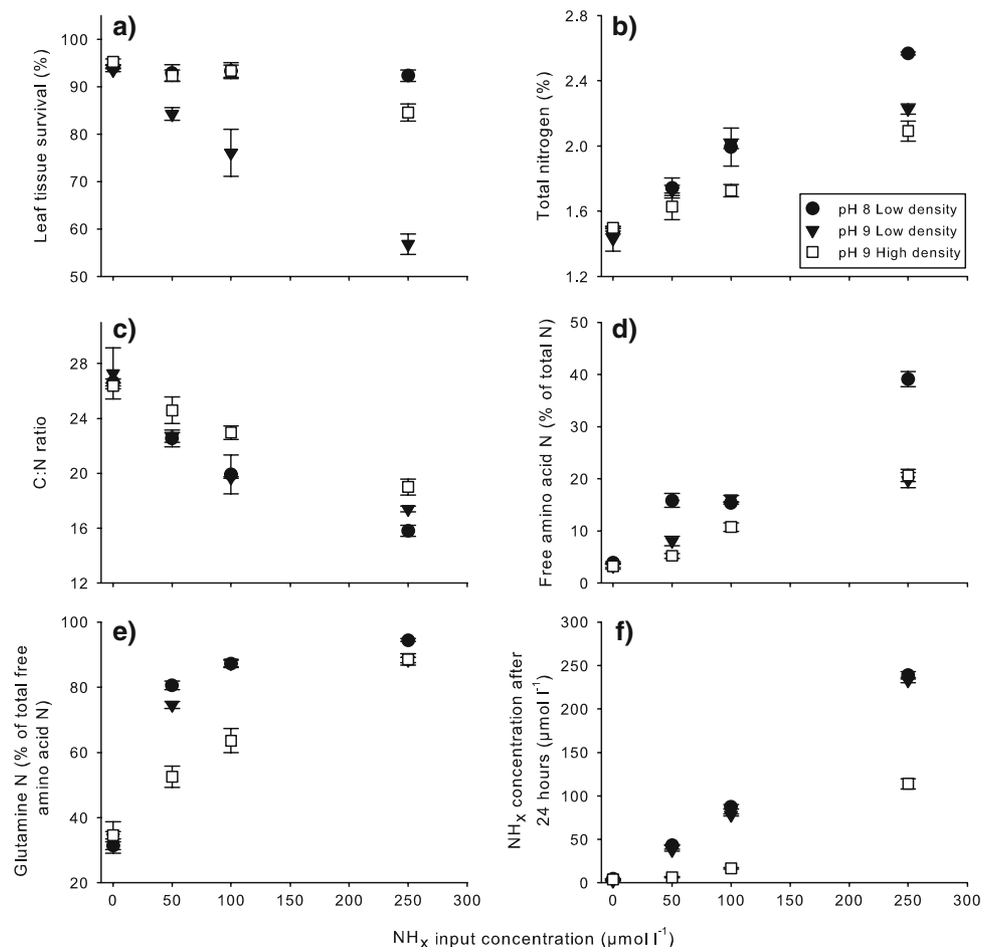


Fig. 2 Eelgrass leaves from **a** 0 μmol l⁻¹ and **b** 250 μmol l⁻¹ NH_x input concentration treatments at low density pH 9 at the end of the short-term experiment. The brown-black discoloration in **b** typifies leaf necrosis

Fig. 3 Effects of the applied NH_x input concentrations (mean \pm SE) in the short-term experiment: **a** leaf tissue survival, **b** total nitrogen (N) content, **c** C:N ratio, **d** free amino acid N relative to total N content, **e** glutamine N relative to total free amino acid N content and **f** NH_x concentration after 24 h



treatments. As in the long-term experiment, excess nitrogen was primarily stored as glutamine. The percentage of glutamine N in the total free amino acid N concentration varied from 33% in the $0 \mu\text{mol l}^{-1}$ treatments to 94% in the $250 \mu\text{mol l}^{-1}$, pH 8 treatment (Fig. 3e).

The NH_x concentration after 24 h was reduced to values around 6 and $17 \mu\text{mol l}^{-1}$ in the high density 50 and $100 \mu\text{mol l}^{-1}$ treatment, respectively. In the $250 \mu\text{mol l}^{-1}$ treatments, NH_x concentrations were still well above $100 \mu\text{mol l}^{-1}$ (Fig. 3f). The reduction of NH_x in the low density treatments was much smaller. The average total uptake ranged from around $10 \mu\text{mol l}^{-1} \text{day}^{-1}$ in the $50 \mu\text{mol l}^{-1}$ treatments to $14 \mu\text{mol l}^{-1} \text{day}^{-1}$ in the $250 \mu\text{mol l}^{-1}$ treatments. NH_x concentrations in the low density pH 9 treatments were slightly but significantly lower compared to their counterparts in the pH 8 treatments.

Discussion

Stress caused by reduced nitrogen in the water layer in seagrasses and other macrophytes has been frequently observed (e.g., Agami et al. 1976; Smolders et al. 1996;

van Katwijk et al. 1997; Bird et al. 1998; Körner et al. 2001; Brun et al. 2002; Cao et al. 2004; Cao et al. 2007; Nimptsch and Pflugmacher 2007). However, the nature of the reported symptoms differ widely. Typical responses are free amino acid accumulation, oxidative stress, inhibited growth, and structural tissue damage. Results from our experiments suggest that differences in response may at least in part be explained by differences in pH of the medium. NH_x toxicity is the sum of the effects caused by NH_4^+ and (gaseous) NH_3 (Nimptsch and Pflugmacher 2007), and the balance between NH_3 and NH_4^+ levels in the NH_x concentration in turn depends on pH (Johansson and Wedborg 1980).

In our first, long-term experiment, conducted at a normal pH level (pH 8.1), structural tissue damage did not appear. However, internal nitrogen concentrations increased linearly with increasing NH_x loads and were much higher than commonly reported for eelgrass. In the $150 \mu\text{M}$ treatment, total N concentrations in the eelgrass leaf tissues were higher than maximum values and nearly 1.5 times higher than average values reported from the field (Duarte 1990). Moreover, free amino acid N concentrations were also much higher than the highest value

reported in literature (Touchette and Burkholder 2002) and consisted mainly of glutamine (around 88%). These results indicate that, like many plants susceptible to NH_4^+ toxicity, eelgrass shoots may not be able to exclude NH_4^+ when exposed to high NH_4^+ levels (Britto et al. 2001; Kronzucker et al. 2001). To prevent accumulation, NH_x then needs to be assimilated by the plant. Glutamine is often the main compound formed in this process, because it is the first amino acid formed in the NH_x -assimilation cycle of plants (Märschner 1995).

In our second, short-term experiment, leaf survival in the low density treatments was strongly affected by the applied NH_x load at pH 9, whereas no effects were visible at pH 8. At first sight, these results may suggest that the observed necrosis should be attributed to increased diffusion of NH_3 into the tissues. After all, the concentration of this species is much higher at pH 9 (28.5% of NH_x) than at pH 8 (3.8% of NH_x) (Johansson and Wedborg 1980). However, this explanation alone does not seem satisfactory, as both the long-term experiment and the pH 8 treatments of the short-term experiment demonstrated that nitrogen also accumulated strongly at pH 8 in eelgrass at the applied NH_x concentrations. It seems likely that disruption of the carbon assimilation (through photosynthesis) rendered the plants susceptible to enhanced external NH_x concentrations at pH 9, as the carbon consuming NH_x assimilation becomes impaired when carbon availability is limited (Märschner 1995). Even though eelgrass is able to utilize bicarbonate (HCO_3^-) next to carbon dioxide (CO_2) for carbon assimilation, carbon may become limited at pH 9 when the total dissolved inorganic carbon (DIC) concentration is at a natural sea water level (± 2.2 mM) (Sand-Jensen and Gordon 1984). This is because with rising pH, CO_2 and HCO_3^- levels drop in favor of CO_3^{2-} , a carbon species which cannot be assimilated. Next, high intracellular NH_x concentrations may cause uncoupling of photophosphorylation by NH_3 in the chloroplasts, further decreasing photosynthesis and related carbon uptake (e.g., Pearson and Stewart 1993; Märschner 1995). This hypothesis is supported by the analyses of the tissue nitrogen compounds in the short-term experiment. Free amino acid and total N concentrations were lower instead of higher in the pH 9 treatments, suggesting that, despite a possible increased intrusion of NH_3 , NH_x assimilation rates were lower at pH 9.

Another important outcome of the short-term experiment is that a high density of eelgrass was able to buffer for NH_x toxicity up to a load of $100 \mu\text{mol l}^{-1}$ NH_x per day, as necrosis only appeared in the highest NH_x treatment. Even though these high density treatments must also have experienced carbon limitation and impaired photosynthesis, survival of the plants was not affected in the three lower NH_x

treatments. Additionally, total tissue nitrogen and free amino acid concentrations were significantly lower in the high density treatments compared to their low density counterparts. This indicates that the potential for NH_x toxicity was alleviated in the high density treatments through the joint uptake of NH_x by the shoots, thereby decreasing ambient NH_x concentrations below the threshold above which NH_x becomes toxic. This view is supported by the measured decrease of NH_x concentrations in the water column within 24 h. The high density treatments were able to lower NH_x concentrations in the 50 and $100 \mu\text{mol l}^{-1}$ treatments to values well below $20 \mu\text{mol l}^{-1}$ within 24 h.

The visual symptoms of NH_x toxicity that appeared in the pH 9 treatments are much alike those reported by Van Katwijk et al. (1997). They also observed a typical brown-black discoloration of the eelgrass leaves within a period of 2 weeks. It is striking, however, that the experimental setup of Van Katwijk et al. was much more similar to the design of our long-term experiment. Both experiments were performed in the same laboratories, lasted 5 weeks, and were performed in glass containers with similar light conditions. Replenishment of the medium was continuous in both experiments, resulting in similar NH_x concentrations in the containers. However, in contrast to the results of Van Katwijk et al., we found no visual symptoms of NH_x toxicity in our long-term experiment. Moreover, the mean total tissue N concentration in our highest treatment was even higher than the highest reported value by Van Katwijk et al. In our opinion, these contradicting results can be best explained by differences in pH. In our experiment, pH was carefully controlled. Van Katwijk et al. mention an average pH of 8.5, which is already somewhat higher than in our experiment (8.0–8.2), but additionally pH was not controlled in their experiment. This lack of pH control most likely allowed for pH fluctuations during the day due to photosynthetic activity. It is therefore not inconceivable that the photosynthetic activity of eelgrass plants may have caused pH spikes of around pH 9 or even higher near the end of each day, leading to carbon limitation and impaired photosynthesis.

Ecological implications

At normal pH levels (i.e., ‘open ocean’ conditions, pH 8.1) and sufficient light for photosynthesis and carbon fixation, acute toxic effects may at least temporarily be prevented by the assimilation of NH_4^+ into free amino acid compounds (van Katwijk et al. 1997; Brun et al. 2002). However, the observed disproportional accumulation of glutamine can be considered an indication of physiological stress due to the elevated NH_4^+ levels in the water layer (Näsholm et al. 1994; Huhn and Schulz 1996; Smolders et al. 2000). Such a chronic exposure to NH_4^+ (i.e., several months or years)

will most likely lead to the classically reported visual symptoms like chlorosis of leaves and the suppression of growth (Britto and Kronzucker 2002). Moreover, phenolic content in the leaves will decrease, due to a changed allocation of carbon skeletons, making the plant more susceptible to pathogens like the “wasting disease”, which destroyed many eelgrass stands in the 1930s (Buchsbbaum et al. 1990; van Katwijk et al. 1997; Vergeer and Develi 1997).

Although high NH_x levels in the water layer at pH 8 most likely cause stress in eelgrass when exposed for longer periods of time, results from our experiments demonstrate that acute toxicity and sudden collapse through mass mortality will probably not occur as long as sufficient light is available. In contrast, elevated NH_x concentrations at enhanced pH levels (i.e., pH 9) can cause severe mortality in eelgrass within days. Thus, mass mortality through NH_x toxicity can be of significant importance in for instance sheltered eutrophicated estuaries, where phytoplankton, macroalgae, and eelgrass itself can cause high pH levels as well as NH_x spikes near the end of the growing season. Additionally, the mechanism may also be important at a normal pH during phytoplankton blooms or along eelgrass depth limits as light availability is poor in those cases.

Interestingly, our results show that the severity of the toxic effect is most likely strongly dependent upon the shoot density of an eelgrass bed. Especially in sheltered stagnant estuaries where mixing between the eelgrass canopy and its surroundings is poor, this could lead to a positive feedback: once there is a high density of eelgrass, its susceptibility to NH_x toxicity becomes lower. If such positive feedback is strong enough, it can lead to alternative stable states in seagrasses (Scheffer et al. 2001; van der Heide et al. 2007). However, in our case, the feedback mechanism may be more complex: a high eelgrass density can also imply a higher photosynthetic activity and thus a higher pH, which in turn may lead to a higher toxicity. Therefore, the outcome in a particular field situation will strongly depend on the NH_x loads, the photosynthetic activity, and the exchange rates between the seagrass meadow and the surrounding water layer (van Nes and Scheffer 2005). The occurrence of alternative stable states may at least in part explain the worldwide observed seemingly high resilience to change, sudden ecosystem collapses, and mass mortalities in seagrasses (van der Heide et al. 2007).

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