



UvA-DARE (Digital Academic Repository)

Low atmospheric nitrogen loads lead to grass encroachment in coastal dunes, but only on acid soils

Remke, E.; Brouwer, E.; Kooijman, A.; Blindow, I.; Roelofs, J.G.M.

DOI

[10.1007/s10021-009-9282-0](https://doi.org/10.1007/s10021-009-9282-0)

Publication date

2009

Document Version

Final published version

Published in

Ecosystems

[Link to publication](#)

Citation for published version (APA):

Remke, E., Brouwer, E., Kooijman, A., Blindow, I., & Roelofs, J. G. M. (2009). Low atmospheric nitrogen loads lead to grass encroachment in coastal dunes, but only on acid soils. *Ecosystems*, 12(7), 1173-1188. <https://doi.org/10.1007/s10021-009-9282-0>

General rights

It is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), other than for strictly personal, individual use, unless the work is under an open content license (like Creative Commons).

Disclaimer/Complaints regulations

If you believe that digital publication of certain material infringes any of your rights or (privacy) interests, please let the Library know, stating your reasons. In case of a legitimate complaint, the Library will make the material inaccessible and/or remove it from the website. Please Ask the Library: <https://uba.uva.nl/en/contact>, or a letter to: Library of the University of Amsterdam, Secretariat, Singel 425, 1012 WP Amsterdam, The Netherlands. You will be contacted as soon as possible.

UvA-DARE is a service provided by the library of the University of Amsterdam (<https://dare.uva.nl>)

Low Atmospheric Nitrogen Loads Lead to Grass Encroachment in Coastal Dunes, but Only on Acid Soils

Eva Remke,^{1,4*} Emiel Brouwer,² Annemieke Kooijman,³ Irmgard Blindow,¹ and Jan G. M. Roelofs⁵

¹Biological Station of Hiddensee, Ernst-Moritz-Arndt-University Greifswald, Biologienweg 15, 18565 Kloster, Germany; ²Research Center B-WARE B.V., Radboud University Nijmegen, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands; ³Institute of Biodiversity and Ecosystem Dynamics, Physical Geography, University of Amsterdam, Nieuwe Achtergracht 166, 1018 WV Amsterdam, The Netherlands; ⁴Bargerveen Foundation, Department of Animal Ecology, Radboud University Nijmegen, Toernooiveld 1, 6525 ED Nijmegen, The Netherlands; ⁵Department of Aquatic Ecology and Environmental Biology, Radboud University Nijmegen, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands

ABSTRACT

The impact of atmospheric N-deposition on succession from open sand to dry, lichen-rich, short grassland, and tall grass vegetation dominated by *Carex arenaria* was surveyed in 19 coastal dune sites along the Baltic Sea. Coastal dunes with acid or slightly calcareous sand reacted differently to atmospheric wet deposition of 5–8 kg N ha⁻¹ y⁻¹. Accelerated acidification, as well as increased growth of *Carex* and accumulation of organic matter, was observed only at acid sites with pH_{NaCl} of the parent material below 6.0. At sites with slightly calcareous parent material, increased N-deposition had no effect. A trigger for grass encroachment seems to be high acidification in early successional stages to below pH_{NaCl} 4.0. Metals like Al or Fe become freely available and may hamper intolerant species. At acid sites, N-mineralization increases

with elevated N-deposition, which may further stimulate *Carex arenaria*. Due to high growth plasticity, efficient resource allocation and tolerance of high metal concentrations, *C. arenaria* is a superior competitor under these conditions and can start to dominate the dune system. *Carex*-dominated vegetation is species-poor. Even at the moderate N-loads in this study, foliose lichens, forbs and grasses were reduced in short grass vegetation at acid sites. Species indicating these first effects of atmospheric deposition on dry, lichen-rich, short grasslands are identified and recommendations for restoration of grass-encroached sites given.

Key words: *Carex arenaria*; nitrogen deposition; species loss; Baltic Sea; acidification; organic matter.

INTRODUCTION

Effects of atmospheric nitrogen deposition on semi-natural ecosystems have been intensively studied in the last three decades. Nutrient-poor and weakly buffered ecosystems are particularly vulnerable. In dry ecosystems in north-western and central

Received 16 April 2009; accepted 10 August 2009;
published online 23 October 2009

*Corresponding author; e-mail: remke@science.ru.nl

Europe, negative impacts of high atmospheric deposition ($15\text{--}30\text{ kg N ha}^{-1}\text{ y}^{-1}$) have been described for forests (Houdijk and others 1993; van Dijk and others 1989, 1990), heathlands (Bobbink and others 1992; Power and others 1998; Roelofs 1986; van den Berg and others 2005) and some types of coastal dunes (Jones and others 2004; Kooijman and others 1998). However, even low levels of atmospheric deposition may have negative effects. For coastal dunes of the Baltic Sea, which only receive $3\text{--}8\text{ kg N ha}^{-1}\text{ y}^{-1}$, critical loads for dry, acid, lichen-rich grasslands as low as $4\text{--}6\text{ kg N ha}^{-1}\text{ y}^{-1}$ have been suggested (Remke and others 2009).

Increased atmospheric N-deposition may affect various ecosystem patterns and processes. Elevated N-loads not only stimulate eutrophication and plant growth, but also lead to acidification and loss of buffer capacity, increasing the availability of toxic metals (Al^{3+} and Fe^{3+}) within the soil (Aerts and Bobbink 1999; Heij and Schneider 1991). Al and Fe concentrations of $50\text{--}100\text{ ppm}$ in plant tissue can harm acid intolerant species and disturb the P-metabolism (Fink 2007). Higher soil Al/Ca-ratios are assumed to be toxic for plants growing at intermediate pH levels (de Graaf and others 1997) as Ca^{2+} inhibits the uptake of Al^{3+} (Marschner 1995).

High N-deposition may also accelerate succession rates in acid grasslands, heathlands and dune pools (Bobbink and others 1998; Achermann and Bobbink 2003). Vegetation becomes dominated by tall graminoids like *Deschampsia flexuosa* (L.) Trin., *Molinia caerulea* (L.) Moench, *Carex arenaria* L. and *Calamagrostis epigejos* (L.) Roth (Bobbink and others 1998; Kooijman and others 1998; Remke and others 2009). Processes behind this species change are a switch from competition for nutrients to a competition for light and space, and resistance of plant species to toxicity such as high metal concentrations or a very low pH (Bobbink and others 1998; de Graaf and others 1997; Kleijn and others 2008). In the Baltic region, dry, lichen-rich dune grasslands are reduced as N-affected dunes become dominated by *Carex arenaria* (Remke and others 2009). Under unpolluted conditions, the succession pathway starts with bare sand, which is then slowly colonized by a few pioneer graminoids like *Corynephorus canescens* P. Beauv. and *Carex arenaria*, by lichens like *Cetraria muricata* (Ach.) Eckfeldt and mosses like *Ceratodon purpureus* (Hedw.) Brid. With time, the bare soil is totally covered by low, but highly characteristic vegetation. These short grasslands contain about $20\text{--}25$ plant species (per relvee of $16\text{--}25\text{ m}^2$), mostly lichens and mosses (Ellenberg 1996). Grey dunes and wet dune slacks

add an essential part to the species diversity of coastal dunes and habitats. Species diversity of coastal habitats comprises $40\text{--}70\%$ of the total diversity in Flanders and The Netherlands (Kooijman 2004; Provoost and Bonte 2004), about 25% on the West Frisian islands (Niedringhaus and others 2008), and forms therefore an essential part of total biodiversity.

Succession from bare sand to fully developed lichen-rich dune grassland may take $50\text{--}70$ years, and is accompanied by a moderate decrease in soil pH ($0.5\text{--}1$ pH units) and accumulation of organic matter (Ellenberg 1996). Eventually taller graminoids, dwarf shrubs or trees become dominant. The sequence from young short via old to tall grassland is one of the major succession lines for Baltic coastal dunes and has been described for the Vistula Spit (Steffen 1931), the Łeba bar (Hueck 1932) as well as the Curonian spit (Paul 1953). However, with increased levels of atmospheric N-deposition, crucial soil processes may change, and lead to dominances of dense, tall grass stands.

The aim of this study is to determine, (i) which soil processes promote the shift from lichen-rich, short, dry grasslands to a dominance of *Carex arenaria*, (ii) which process changes can be linked to atmospheric N-deposition, (iii) to specify characteristics of *Carex arenaria* stands under different atmospheric N-loads, and (iv) to describe the effects of elevated deposition loads on species composition and richness. Finally, early indicators for the influence of low atmospheric N-deposition levels are listed, and recommendations for restoration of dry, coastal dunes dominated by tall graminoids are given.

Field Sites

The 19 coastal dune sites are spread over a large geographical gradient from the Kattegat in the west of the Southern Baltic to the Baltic proper in the east (Figure 1; Remke and others 2009). The research area therefore stretches from the Atlantic through Central Europe to the Baltic terrestrial ecoregion (Olson and others 2004). The average annual temperature decreases from 8°C in Denmark to 5°C in Estonia (Table 1) as the climate gets more continental. Another gradient exists for sea water salinity and therefore salt-spray. Salinity decreases steeply within in the Kattegat from 24 to approximately 10 PSU and ranges mainly between 7 and 8 PSU in the Southern Baltic and Baltic proper (Table 1). All 19 sites have parent sand material with pH_{NaCl} below 6.5 and receive precipitation of $500\text{--}700\text{ mm y}^{-1}$ (Table 1). All sites are open dunes



Figure 1. Map of field sites (reprinted from Remke and others 2009, with permission from Elsevier).

Table 1. Field Sites with Synonym, Country, Geographical UTM Coordinates (from Google Earth), Long-Term Annual Mean Values for Temperature and Precipitation, and Sea Water Salinity

Name/synonym	Country	Longitude, UTM	Latitude, UTM	Temperature, °C	Precipitation, mm y ⁻¹	Salinity, PSU
Akmensrags ¹	Latvia	56411065	20989034	6.8	627	7.2
Dünenheide/Hiddensee ²	Germany	54550269	13099371	8.0	564	9.0
Gellen/Hiddensee ²	Germany	54479486	13064725	8.0	564	9.0
Harilaid ²	Estonia	58489564	21843912	5.6	703	6.9
Holtemmen/Læsø ⁵	Denmark	57302031	10994222	7.9	576	24.0
Keibu ⁴	Estonia	59241349	23739389	5.1	686	7.0
Korshage ⁶	Denmark	55974190	11777358	7.3	500	18.4
Nagliu/Curonian Spit NP ⁷	Lithuania	59659837	21083870	7.8	750	7.3
Nida/Curonian Spit NP ⁷	Lithuania	55284024	20958343	7.8	750	7.3
Pajuris ⁸	Lithuania	55843364	21062018	7.0	735	7.3
Pape ¹	Latvia	56150317	21027620	6.8	627	7.2
Pärisea ⁴	Estonia	59659837	25683665	5.1	686	6.4
Pavilosta ¹	Latvia	56893958	21191670	6.8	627	7.0
Raghhammer/Bornholm ⁹	Denmark	55015227	14926422	7.7	572	7.9
Leba/Slovinski NP ¹⁰	Poland	54682510	17101320	7.7	575	7.5
Sandhammeren ³	Sweden	55378837	14180429	7.4	600	7.7
Syrodde/Læsø ⁵	Denmark	57323485	11195125	7.9	576	24.0
Tönnersa ³	Sweden	56555071	12947505	7.3	700	18.1
Torsö ³	Sweden	55999030	14657380	7.4	550	7.7

NP = national park.

Sources for climate data: ¹ www.worldclimate.com, station Liepaja; ² www.worldclimate.com, station Greifswald; ³ temperature: Eggertsson-Karlström (2004); precipitation: Raab and Vedin (1995); ⁴ Ratas and Nilson (1997); ⁵ Walter and Lieth (1967) in Biermann (1999); ⁶ temperature: www.dmi.dk, regions København and Nordsjælland; precipitation: Jensen (1986); ⁷ Anonymous (2004); ⁸ www.wetter.com, station Kleipeda; ⁹ www.worldclimate.com, station Duoedde; ¹⁰ temperature: www.worldclimate.com, station Leba; precipitation: Walna and others (2003). Salinity values are obtained from the mean grid cell values for the period 1900–2005 (Feistel and others 2008).

without any physical barrier like a forest towards the sea, and have less than 10% cover of trees. During the last decade, management impacts such as grazing or burning have not been intensive.

The dataset was separated into two main groups, with pH of the parent material below ('acid') or above ('slightly calcareous') 6.0 (Table 2), that is, below or within the carbonate buffer range. Above pH 6.5–6.2, the system is buffered by dissolution of calcium carbonate below pH 6.0 by cation exchange, and below pH 5.0 by aluminium and iron buffer systems (Scheffer and Schachtschabel 1998). In addition, both pH-groups were separated into two N-deposition classes below and above a total wet N-deposition of $5 \text{ kg N ha}^{-1} \text{ y}^{-1}$, estimated by N-content of the biomonitor *Cladonia portentosa* (Remke and others 2009; Table 2). The calculated total wet N-deposition ranged from 2.6 to 7.8 total N $\text{kg ha}^{-1} \text{ y}^{-1}$, but critical loads were estimated as 4–6 $\text{kg N ha}^{-1} \text{ y}^{-1}$ (Remke and others 2009). The N-content in *C. portentosa* proved to be a suitable biomonitor of N-deposition in remote areas as it explained approximately 50% of the variation in wet N-deposition measured at nearby EMEP stations (regression equation: *C. portentosa* tissue N [%] = $0.0228 \times \text{N-deposition} [\text{kg ha}^{-1} \text{ y}^{-1}] + 0.3385$, $R^2 = 0.5223$, $P = 0.008$; further discussion see Remke and others 2009).

MATERIALS AND METHODS

In each of the 19 dune sites, three $2 \times 2 \text{ m}^2$ plots were selected in June–July 2005, in a sequence from young short via old short to tall grassland. First, early successional stage is characterized by short, dry grassland, with at least 30% open sand area, second stage by older and lichen-rich, short grassland, with open sand below 5%, and third stage is dominated by tall grasses with at least approximately 50% cover of sand sedge (*Carex arenaria*). In each plot, species cover was estimated. If necessary, species were dried and identified using a microscope and coloring techniques. For species determination the following literature was used: for vascular plants Rothmaler and others (2002), Oberdorfer (1994) and Hegi and Conert (1998), additionally for *Salix* spp. Lautenschlager-Fleury and Lautenschlager (1994), for mosses Frahm and Frey (1992) and for lichens Wirth (1995). Within each plot, aboveground biomass was sampled in a circle with a diameter of 29.5 cm. Within each circle, thickness of the humus horizon (Ol, Of, Oh) was measured (Finnern and others 1994). Below this, a soil sample mixed out of three soil cores (each core cutter 100 cm^3) was collected from the top 10 cm of the mineral soil layer. In addition, at each site, three soil samples (top 2–3 cm) were

Table 2. Classification of Field Sites

Field site	Parent sand, pH	pH, class	Lichen-N, %		Calc. total N, $\text{kg ha}^{-1} \text{ y}^{-1}$	Deposition
			Mean	SE		
Keibu	5.64	Acid	0.409	0.020	2.57	Low
Pärispea	5.82	Acid	0.425	0.016	3.12	Low
Syrodde	5.30	Acid	0.451	0.017	4.02	Low
Sandhammaren	5.27	Acid	0.463	0.015	4.39	Low
Korshage	5.43	Acid	0.497	0.026	5.54	High
Dunenheide	5.63	Acid	0.522	0.019	6.39	High
Torsö	5.09	Acid	0.537	0.020	6.89	High
Gellen	5.02	Acid	0.545	0.017	7.19	High
Tönnersa	5.50	Acid	0.554	0.032	7.48	High
Nagliu	6.82	Slightly calcareous	0.404	0.018	2.41	Low
Akmensrags	6.98	Slightly calcareous	0.428	0.025	3.23	Low
Nidda	6.87	Slightly calcareous	0.439	0.022	3.60	Low
Pavilosta	6.85	Slightly calcareous	0.448	0.016	3.91	Low
Harilaid	6.59	Slightly calcareous	0.484	0.017	5.10	High
Holtemmen	6.15	Slightly calcareous	0.496	0.019	5.53	High
Leba	6.79	Slightly calcareous	0.504	0.023	5.81	High
Pajuris	6.42	Slightly calcareous	0.505	0.015	5.82	High
Raghammer	6.63	Slightly calcareous	0.513	0.013	6.10	High
Pape	6.54	Slightly calcareous	0.562	0.017	7.76	High

Sites with pH_{NaCl} above 6.0 of the parent sand material are classified as 'slightly calcareous' sites with pH_{NaCl} below 6.0 as 'acid'. Sites with calculated total wet N-deposition below $5 \text{ kg N ha}^{-1} \text{ y}^{-1}$ are classified as N-unaaffected or 'low' N-deposition sites, sites with calculated total wet N-deposition above $5 \text{ kg N ha}^{-1} \text{ y}^{-1}$ as N-affected or 'high' N-deposition sites. Total deposition of N was calculated using the lichen biomonitor (calc. total N; method see Remke and others 2009).

taken along the first dune ridge towards the open sea to determine the pH of the parent sand material. All samples were dried at 70°C (plant) and 40°C (soil) for 24 h.

The lichen, *Cladonia portentosa* (Dufour) Coem., was used as biomonitor for atmospheric N deposition (Remke and others 2009). Within each plot, ten 10 × 10 cm² samples of *C. portentosa* were cut with a pair of scissors out of the center of a lichen patch of approximately 0.4–0.5 m diameter (Søchting 1995). Only the top 2 cm of the *Cladonia* were collected to exclude nutrient uptake from the soil. The samples were cleaned roughly and stored in a paper bag. In the laboratory, all samples were cleaned thoroughly, washed briefly in distilled water and dried at 70°C for 24 h.

Fresh biomass was separated into cryptogams (lichens and mosses), vascular plants and litter, and weighed after drying at 70°C for 24 h. A subsample of sand sedge was taken for further analysis. Sand sedge and *Cladonia portentosa* samples were ground in a centrifugal mill (rotational speed 18,000 for 1–2 × min, FRITSCH pulverisette 14, Idar-Oberstein, Germany). Total nitrogen and CN-ratio of plant and soil material (finely ground in a centrifugal ball mill, Fritsch, Idar-Oberstein, Germany) were determined with a C/N-analyzer (CHNOS element analyzer vario EL III, elemental Analysensysteme, Hanau, Germany). Total P was analyzed for sand sedge only, with 200 mg ground material digested in sealed Teflon vessels in a Milestone microwave oven (type Ethos D, Milestone Inc., Sorisole, Italy) after addition of 4 ml HNO₃ (65%) and 1 ml H₂O₂ (30%) (Kingston and Haswell 1997).

Soil samples were sieved with a 2 mm mesh-size before the following analyses. Soil organic matter content was determined as loss on ignition (LOI) at 550°C for 8 h, and pH was measured in 0.2 M NaCl. Total extractable amounts of Al, Ca, Fe, Mg, Mn, P, S, and Zn were measured in these 0.2 M NaCl extracts, and Na in a double deionized water extract with inductively coupled plasma emission spectrophotometry (ICP-OES: IRIS Intrepid II XDL, Thermo Fisher Electric, Breda, The Netherlands). Nitrogen mineralization was measured in a laboratory incubation experiment of 26 days, with 60 g of sieved soil in glass beakers sealed with parafilm kept at 40% water holding capacity and 25°C in total darkness. NO₃⁻ and NH₄⁺ were measured at the start and end of the incubation period, using extractions with double deionized water and 0.2 M NaCl, respectively. *Ortho*-P was measured in double deionized water extracts at the start of the incubation period. Nitrate, ammonium and phosphate were measured

colorimetrically with an Auto Analyzer 3 system (Bran + Luebbe, Norderstedt, Germany), using ammonium molybdate (Henriksen 1965), hydrazine sulphate (Technicon 1969) and salicylate (Grasshoff and Johannsen 1977), respectively.

Linear models and linear mixed effect models were fitted using R (R Development Core Team 2008), followed by model justification procedures. Mixed effect models were applied to overcome spatial pseudo-replication within the dataset (field site as random factor). If fitted models were not justified, generalized linear models (glm) with gamma error distributions were fitted or data were transformed by log or (double) square root before regression analysis. Multiple regressions were carried out starting with all environmental factors and subsequently simplified using stepwise backward deletion until the minimal adequate model was reached. To avoid over-dispersion no more variables than the replicate number divided by three were fitted (Crawley 2005, 2007). Tests between two classes were performed by Student's *t* test, if the data were normally distributed, otherwise the non-parametric Kruskal–Wallis test was performed.

RESULTS

Acidification

Acidification rates differed between early and later successional stages. Initial acidification, that is, differences in pH between parent material and the first succession stage, was negatively correlated with acidification during subsequent succession (Figure 2). At sites with small pH differences between parent material and the first stage of succession, the pH decreased by nearly two units during the succession from the first to the third stage. Sites following this pattern were Akmensrags, Nagliu, Pajuris, Syrodde, Pape and Gellen (Figure 2). Sites with high pH differences (~2 units) between parent material and initial succession stage, showed minimal pH decrease with further succession. Dünenheide, Harilaid, Pavilosta, Holtemmen, Keibu and Korshage were examples for this pattern. The early acidification, that is, the high pH-decrease between the parent sand material and the first stage, shows no link to N-deposition levels. The early decrease in pH (parent sand—first stage) cannot be linked to N-deposition (regression results: acid *P* = 0.595; slightly calcareous *P* = 0.845; all *P* = 0.660). Only the pH itself differed with N-deposition. Especially at acid sites and later successional stages the pH was significantly lower (0.3–0.5 units) at N-affected sites (Table 3).

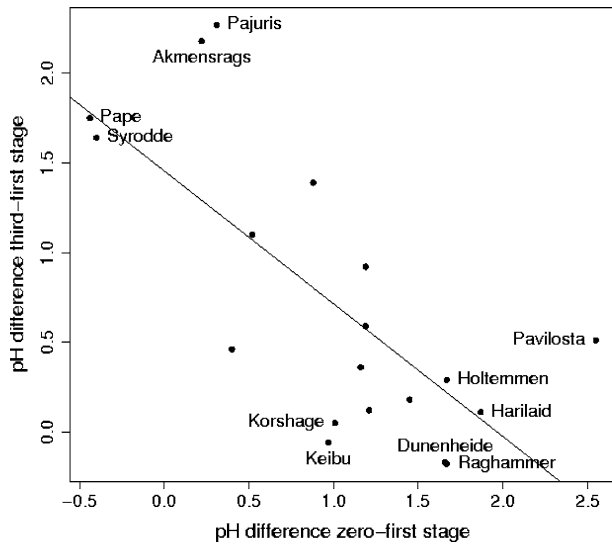


Figure 2. Relationship between the pH difference of the parent sand material and the first vegetation succession stage and the pH difference within succession, that is, first and the third succession stage. Every point is a site and extreme sites are labelled. For example, Pajuris or Akmensrags have a low pH decrease between the parent sand material and the first succession stage, and Pavilosta and Harilaid have a high pH difference within later succession (regression equation: $y = -0.74x + 1.45$, $R^2 = 0.5166$, $P = 0.001$).

Eutrophication

Soil pH and organic matter (LOI) were closely related and changed during succession, but were also affected by N-deposition (Figure 3). Organic matter increased exponentially with decreasing pH, but only in areas with high N-deposition. Plots with more than 1.0–1.5% soil organic matter and a pH below 4.0 belonged mainly to the third succession stage at N-affected sites, which are densely covered by *Carex arenaria*. A significant increase in organic matter with N-deposition occurred only at sites with acid parent material (Table 3). At these sites, organic matter concentration was two to four times higher at N-affected than at N-unaffected sites. At sites with slightly calcareous parent material, however, organic matter increased during succession, but did not differ between low and high N-deposition areas (Table 3).

During succession, net N-mineralization generally increased from the first to the third stage (Table 3). The amount of N-mineralization per unit soil organic matter was positively correlated with pH (Figure 4) and furthermore related to N-deposition. The increase of N-mineralization/LOI with increasing pH was three times steeper at N-affected than at N-unaffected sites (Figure 4), and N-min-

eralization in total was elevated two to three times at sites with high N deposition (Table 3).

Toxicity

The Al/Ca-ratio increased with increasing N-deposition only at sites with acid parent material (second stage, Table 4), and was twice as high at N-affected than at N-unaffected sites (second stage, Table 3). At acid sites, the Al/Ca-ratio exceeded the values of one in all succession stages. At sites with slightly calcareous parent material, Al/Ca-ratios were 5–10 times lower. Salt-extractable Fe soil concentrations were at least two times higher at acid than at slightly calcareous sites. At acid sites, the Fe content was 1.5–3 times higher under high compared to low N-deposition.

Characteristics of *Carex arenaria* Plants and Vegetation Stands

Increased atmospheric N-deposition (measured as lichen-N) was positively correlated to the N-content of *Carex* tissue, but only in early succession stages at acid sites and all sites pooled (Table 4). N/P-ratios (*Carex*) were positively correlated to N-deposition in all three succession stages at acid sites (Table 4). Tissue P-concentrations of *Carex arenaria* did not differ among sites or succession stages, and ranged between 51.2 and 56.6 $\mu\text{mol P g}^{-1}$. There was no significant relationship between tissue-N of *Carex* and total N, NH_4^+ or NO_3^- in the soil.

At acid sites, cover of *Carex arenaria* significantly increased under N-affected conditions in later stages of succession (Tables 3 and 4, Figure 5). *Carex* cover was 1.5–2 times higher under N-affected conditions. In early stages, or slightly calcareous sites, *Carex* cover was not affected by N-deposition. In the third succession stage, *Carex* cover was negatively correlated to species numbers of lichens and mosses ($y = -0.09x + 8.48$, $P = 0.000$) and positively to total biomass of vascular plants ($y = 0.47x + 33.70$, $P = 0.027$). *Carex* cover was not significantly correlated to soil organic matter. Furthermore, longitude, latitude, climate data (temperature, precipitation) and salinity were not significantly correlated to *Carex* cover in multiple regression.

Species Richness

At acid sites, the species number of foliose lichens, as well as number of all lichen species together (second and third stage; Figure 6), and grasses and forbs (third stage) were significantly lower (two to three species) under high N-deposition (Table 5).

Table 3. Soil and Plant Data for All Sites, Separated for pH Classes (Acid, Slightly Calcareous), Deposition Classes (N-Unaffected and N-Affected) and Vegetation Succession Stages (First, Second and Third stage)

Deposition	N-unaffected			N-affected		
	First	Second	Third	First	Second	Third
Acid sites						
Lichen-N, %	0.428 (0.06)	0.437(0.06)	0.463 (0.00)	0.541 (0.03)	0.531 (0.06)	0.533 (0.05)
LOI, %	0.28 (0.03)	0.50 (0.05)	0.41 (0.03)	0.38 (0.07)	1.15 (0.19)	1.46 (0.26)
Ol, cm	0.1 (0.04)	0.3 (0.05)	0.4 (0.10)	0.0 (0.00)	0.10 (0.04)	0.8 (0.10)
pH, NaCl	4.91 (0.31)	4.25 (0.11)	3.94 (0.02)	4.27 (0.08)	3.74 (0.06)	3.67 (0.09)
C/N-ratio soil	10.9 (1.2)	11.6 (0.6)	11.8 (1.6)	8.7 (0.9)	14.2 (0.7)	13.5 (0.9)
N-miner., g/m ³	0.69 (0.12)	0.84 (0.12)	0.87 (0.19)	1.27 (0.16)	1.69 (0.21)	2.65 (0.36)
N-NH ₄ , g/m ³	0.56 (0.11)	0.78 (0.11)	0.81 (0.22)	0.91 (0.13)	0.93 (0.06)	1.22 (0.12)
N-NO ₃ , g/m ³	1.20 (0.30)	2.29 (0.70)	0.99 (0.28)	2.41 (0.63)	3.84 (0.86)	5.41 (1.08)
NH ₄ /NO ₃ ratio	0.57 (0.1)	0.74 (0.2)	0.92 (0.3)	0.54 (0.1)	0.49 (0.1)	0.36 (0.1)
Al/Ca-ratio soil	1.27 (0.41)	0.40 (0.13)	0.42 (0.1)	1.19 (0.2)	1.04 (0.2)	0.98 (0.39)
Fe, µmol/kg	9.03 (1.96)	20.56 (3.26)	40.28 (2.12)	13.00 (1.97)	65.73 (11.32)	74.42 (14.54)
CEC, µmol/kg	1190 (504)	1276 (270)	662 (106)	624(142)	1250 (431)	1781 (317)
<i>Ortho</i> -P, µmol/kg	9.98 (2.05)	24.70 (5.14)	42.49 (5.08)	27.53 (7.17)	31.22 (8.13)	63.18 (8.69)
N/P-ratio <i>Carex</i>	7.8 (1.1)	7.1 (0.4)	8.1 (0.1)	11.5 (0.5)	9.6 (0.5)	9.6 (0.5)
N <i>Carex</i> , %	11.43 (0.54)	11.72 (0.99)	12.95 (1.05)	17.48 (1.44)	14.61 (0.54)	15.29 (0.63)
P <i>Carex</i> , %	1.71 (0.35)	1.67 (0.12)	1.62 (0.15)	1.59 (0.18)	1.56 (0.09)	1.62 (0.08)
<i>Carex</i> cover per plot, %	2.9 (1.1)	6.2 (2.0)	47.7 (10.2)	5.6 (1.4)	12.3 (2.1)	77.1 (4.7)
Vascular plant biomass, g/m ²	66.8 (14.1)	219.4 (63.4)	954.8 (107.8)	78.7 (10.4)	539.5 (164.6)	1066.4 (94.9)
Lichen and moss biomass, g/m ²	16.7 (11.1)	768.3 (120.4)	51.8 (32.9)	13.1 (5.6)	868.3(173.3)	371.2 (170.7)
Slightly calcareous sites						
Lichen-N, %	0.441 (0.07)	0.442 (0.08)	0.441 (0.07)	0.516 (0.06)	0.516 (0.06)	0.519 (0.08)
LOI, %	0.31 (0.03)	0.74 (0.11)	0.89 (0.07)	0.26 (0.05)	0.64 (0.08)	0.84 (0.14)
Ol, cm	0.05 (0.02)	0.80 (0.18)	1.28 (0.22)	0.07 (0.04)	0.59 (0.11)	1.25 (0.23)
pH, NaCl	5.58 (0.26)	4.90 (0.24)	4.73 (0.2)	5.70 (0.3)	4.74 (0.2)	4.57 (0.21)
C/N-ratio soil	16.0 (3.4)	12.0 (0.7)	12.0 (0.62)	12.0 (0.87)	12.7 (0.93)	13.5 (0.72)
N-miner., g/m ³	1.19 (0.11)	1.38 (0.36)	3.00 (0.37)	1.10 (0.15)	1.81 (0.27)	3.33 (0.64)
N-NH ₄ , g/m ³	0.35 (0.08)	0.65 (0.12)	1.05 (0.09)	0.42 (0.06)	0.52 (0.11)	0.90 (0.17)
N-NO ₃ , g/m ³	1.19 (1.08)	5.17 (1.30)	4.08 (0.88)	1.02 (0.69)	4.24 (0.63)	3.20 (1.33)
NH ₄ /NO ₃ ratio	0.25 (0.1)	2.00 (0.8)	0.35 (0.05)	0.16 (0.03)	1.02 (0.44)	0.42 (0.14)
Al/Ca-ratio soil	0.74 (0.4)	0.12 (0.1)	0.71 (0.6)	0.07 (0.03)	0.19 (0.11)	0.08 (0.02)
Fe, µmol/kg	4.86 (0.66)	13.24 (2.69)	24.81 (5.64)	5.46 (1.08)	18.32 (4.79)	22.04 (4.81)
CEC, µmol/kg	1353 (224)	2600 (342)	2663 (342)	1141 (127)	1915 (274)	2356 (320)
<i>Ortho</i> -P, µmol/kg	12.20 (1.60)	49.68 (13.57)	54.52 (9.84)	22.78 (7.37)	64.37 (12.71)	70.39 (14.83)
N/P-ratio <i>Carex</i>	9.9 (0.7)	7.9 (0.5)	7.9 (0.4)	9.3 (0.6)	7.8 (0.4)	7.0 (0.28)
N <i>Carex</i> , %	13.02 (0.42)	12.41 (0.46)	13.27 (0.77)	14.91 (0.65)	12.76 (0.38)	13.71 (0.61)
P <i>Carex</i> , %	1.40 (0.09)	1.62 (0.08)	1.73 (0.13)	1.76 (0.20)	1.73 (0.09)	1.99 (0.12)
<i>Carex</i> cover per plot, %	4.3 (0.7)	5.8 (0.8)	60.6 (2.5)	4.9 (1.1)	7.0 (1.2)	65.1 (3.29)
Vascular plant biomass, g/m ²	86.3 (9.0)	233.7 (41.0)	905.0(68.0)	84.4(10.2)	238.5 (30.4)	897.9 (78.7)
Lichen and moss biomass, g/m ²	18.6 (11.6)	893.0 (80.6)	180.8(72.0)	29.3(19.0)	844.1 (97.3)	147.1 (68.5)

LOI = loss on ignition, Ol = thickness of litter layer, CEC = cation exchange capacity.

Values are arithmetic means (\pm 1SE). Significant differences ($P < 0.05$) between deposition classes (N-affected and N-unaffected) are shown in bold.

Foliose lichens such as *Cladonia glauca* Flörke, *Cl. macilenta* (Leighton) Arnold, *Cl. coccifera* (L.) Willd. and *Cetraria aculeata* (Schreber) Fr./*muricata* (Ach.) Eckfeldt, grasses such as *Festuca polesica* Zapał. and *Ammophila arenaria* (L.) Link, and forbs such as *Hypochaeris radicata* L., *Thymus serpyllum* L. and *Hieracium umbellatum* L. were absent or had lower cover at N-affected sites. In contrast, *Carex arenaria*

and *Corynephorus canescens* were more abundant at acid sites with high N-deposition, and *Rumex acetosella* L. was exclusively found at these sites.

The most important predictors for the decrease in species numbers at acid sites were selected with multiple regression. For the second succession stage, the model explained 20–50% of the variance, with Al/Ca-ratio and its (statistical) interaction

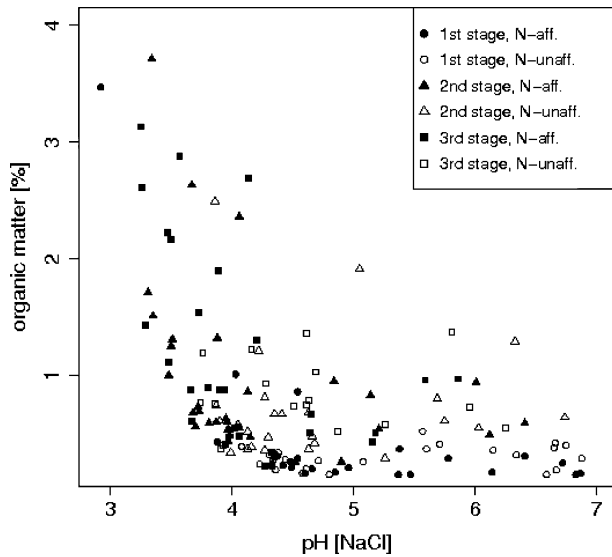


Figure 3. Organic matter content (LOI) versus pH of all vegetation units.

with pH and N-deposition as most important factors. For the third stage, only N-deposition was a significant predictor for species richness and explained 30–60% of the variance (Table 6). Species with preference for low Al/Ca-ratios were *Hieracium umbellatum* (higher coverage), *Cladonia glauca* (higher coverage and occurrence), and *Jasione montana* L., *Cladonia coccifera* and *Cladonia florkeana* (Fr.) V.Wirth (higher occurrence). Species excluded from sites with high Al/Ca-ratios were *Cladonia glauca*, *Viola* ssp. and *Luzula* ssp. However, *Rumex acetosella*, *Anthoxanthum odoratum* L. and *Corynephorus canescens* were more abundant at high Al/Ca-ratios.

At sites with slightly calcareous parent material, the effect of N-deposition was less pronounced than at acid sites, and even slightly positive for overall species richness (Table 5). In multiple regression, Al/Ca-ratio and pH mainly determined species richness (Table 6). N-affected sites had one species of foliose lichen (third stage), grass (second stage) and forb (first stage) more, but one moss species less (second stage). *Cladonia coniocrea* auct. (foliose lichen), *Hypochaeris radicata* (forb), *Anthoxanthum odoratum* and *Festuca rubra* L. (grasses) were more abundant at high N-deposition. *Dicranum scoparium* Hedw. (second stage) had exceptionally high coverages (30–40%) at high N-deposition. However, *Pleurozium schreberi* (Brid.) Mitt. (moss), *Cetraria aculeata/muricata* and *C. islandica* (L.) Ach. (first and second stage) occurred only at N-unaffected sites.

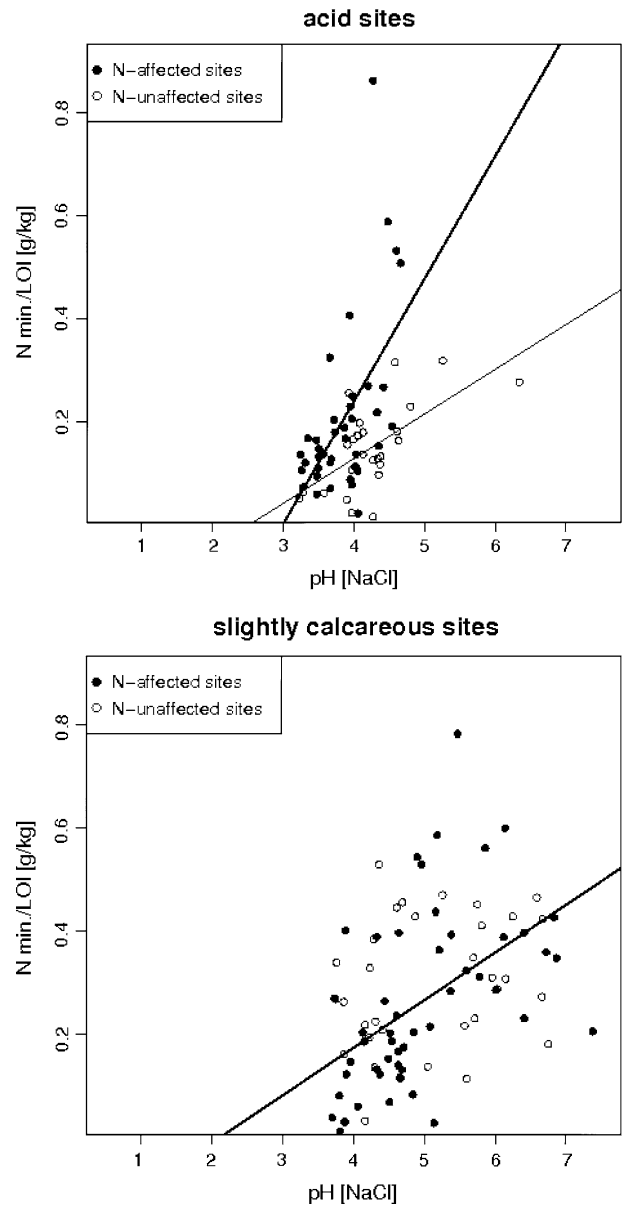


Figure 4. N-mineralization per day per LOI (g kg^{-1}) against the pH (NaCl) for acid and slightly calcareous sites separately. (Regression results: acid sites: N-affected $y = 0.238x - 0.712$ ($P = 0.000$), N-unaffected $y = 0.087x - 0.218$ ($P = 0.001$); slightly calcareous sites: N-affected $y = 0.092x - 0.194$ ($P = 0.005$), N-unaffected not regression equation as $P = 0.104$).

DISCUSSION

Which Processes Change During Grass Encroachment?

At these 19 coastal dune sites within the Baltic Sea region even medium loads of wet N-deposition ($5\text{--}8 \text{ kg N ha}^{-1} \text{ y}^{-1}$) have an impact on soil and

Table 4. Regression Results of Various Factors (Dependent Factor) with Atmospheric N-Deposition (Measured as Lichen-N; Independent Factor)

Depending factor	Acid sites		Slightly calcareous	All sites	
PH-difference parent sand—First stage		$P = 0.595$	$P = 0.845$		$P = 0.660$
Al/Ca-ratio (second stage)	$y = 0.85x - 3.34$	$P = 0.034$	$P = 0.807$		$P = 0.179$
Carex tissue N (first stage)		$P = 0.137$	$P = 0.196$	$y = 3.45x - 2.10$	$P = 0.021$
Carex tissue N (second stage)	$y = 2.79x - 0.73,$	$P = 0.040$	$P = 0.446$	$y = 1.74x + 4.35$	$P = 0.013$
Carex tissue N/P-ratio (first stage)	$y = 3.86x - 8.95,$	$P = 0.013$	$P = 0.919$	$y = 1.93x - 0.48$	$P = 0.058$
Carex tissue N/P-ratio (second stage)	$y = 2.53x - 4.10,$	$P = 0.012$	$P = 0.331$	$y = 1.83x - 0.84$	$P = 0.009$
Carex tissue N/P-ratio (third stage)	$y = 3.23x - 7.53,$	$P = 0.047$	$P = 0.778$		$P = 0.122$
Carex cover [%] (third stage)	$y = 7.90x - 29.07,$	$P = 0.039$	$P = 0.806$	$y = 4.72x - 14.92$	$P = 0.040$

Climatic factors (temperature, precipitation), salinity and UTM-coordinates had no significant impact in multiple regression. Regression results of the listed factors were not significant in other successional stages and are not shown.

vegetation factors. At these medium N-loads factors are changing significantly only at acid sites (parent sand pH < 6.0) not at slightly calcareous sites (parent sand pH > 6.0). The pH of the parent sand material is a main factor responsible for the reaction of the system to these low to medium N-loads. Although pH-differences are small between acid and slightly calcareous sites, they determine which buffer range is acting and how quickly the buffering capacity of cations is depleted, if toxic levels of soluble metals can occur or if mineralization processes are hampered.

The acidity of the soil has a major influence on soil processes. Some field sites show the expected pattern of acidification during vegetation succession in coastal dunes. The pH remains almost constant during the transition from parent sand material to first stage, but decreases with increasing successional age (Ellenberg 1996). In contrast to these are sites where the pH decreases already between the parent sand material and the first succession stage, after which it remains stable. This pH reduction during the early successional stages was not correlated to N deposition, but may instead be caused by anthropogenic sulphur deposition, for example, ship traffic. Sulphur generally generates two times more protons than reduced or oxidized nitrogen (NEG-TAP 2001). Pure chemical reactions might prevail in the pure sands at initial successional stages, whereas biological transformations of N such as nitrification may dominate the acidifying processes in later succession stages with higher soil organic matter contents. This is supported by the lower pH values of later successional stages at higher N-deposition in this study.

As soil pH decreases, organic matter (LOI) accumulates exponentially (this paper; Stützer 1998; Scheffer and Schachtschabel 1998). In these dry,

oligotrophic systems, organic matter retains moisture and stabilizes the soil micro-climate. Most typical dry dune plant species are adapted to fluctuating, extreme temperatures, drought and frequent soil moisture changes. The more stabilized conditions, which prevail during organic matter accumulation, support the growth of ruderal, nitrophilous species at the cost of typical dry dune species like *Jasione montana*. Graminoids such as *Carex arenaria*, *Calamagrostis epigejos* and *Festuca rubra* become dominant, where a sufficient layer of organic matter is available or the top sand layer is nutrient rich (Boorman and Van der Maarel 1997). This observation is supported by the current dataset. At acid sites, the total standing vascular plant biomass and the cover of *C. arenaria* are 1.5–3 times higher at N-affected than at N-unaffected sites.

At older successional stages, the higher N-content and N/P-ratio in *Carex* tissue at N-affected sites is in contrast with the higher C/N-ratio in the soil. Different explanations are possible. One is that surplus mineralized N is taken up directly by *Carex*. In addition, in acid soils more N might be available for vascular plants in competition with microbes as microbial communities seem to have a lower demand for N (Kooijman and Besse 2002; Kooijman and others 2008). Another possible explanation is, that NO_3^- , which is not taken up, is readily leached. Furthermore, the immobilized N can be bound in structures with higher C/N-ratios like fungi in these acid soils, which might then result in a higher C/N-ratio for the total soil.

Although nitrification is hampered at a pH below 4.2 (Aerts and Bobbink 1999; Roelofs and others 1985), the higher amount of organic matter at acid, N-affected sites nevertheless increases N-mineralization to the same level as at slightly calcareous sites (Table 3). At acid sites, microbes have

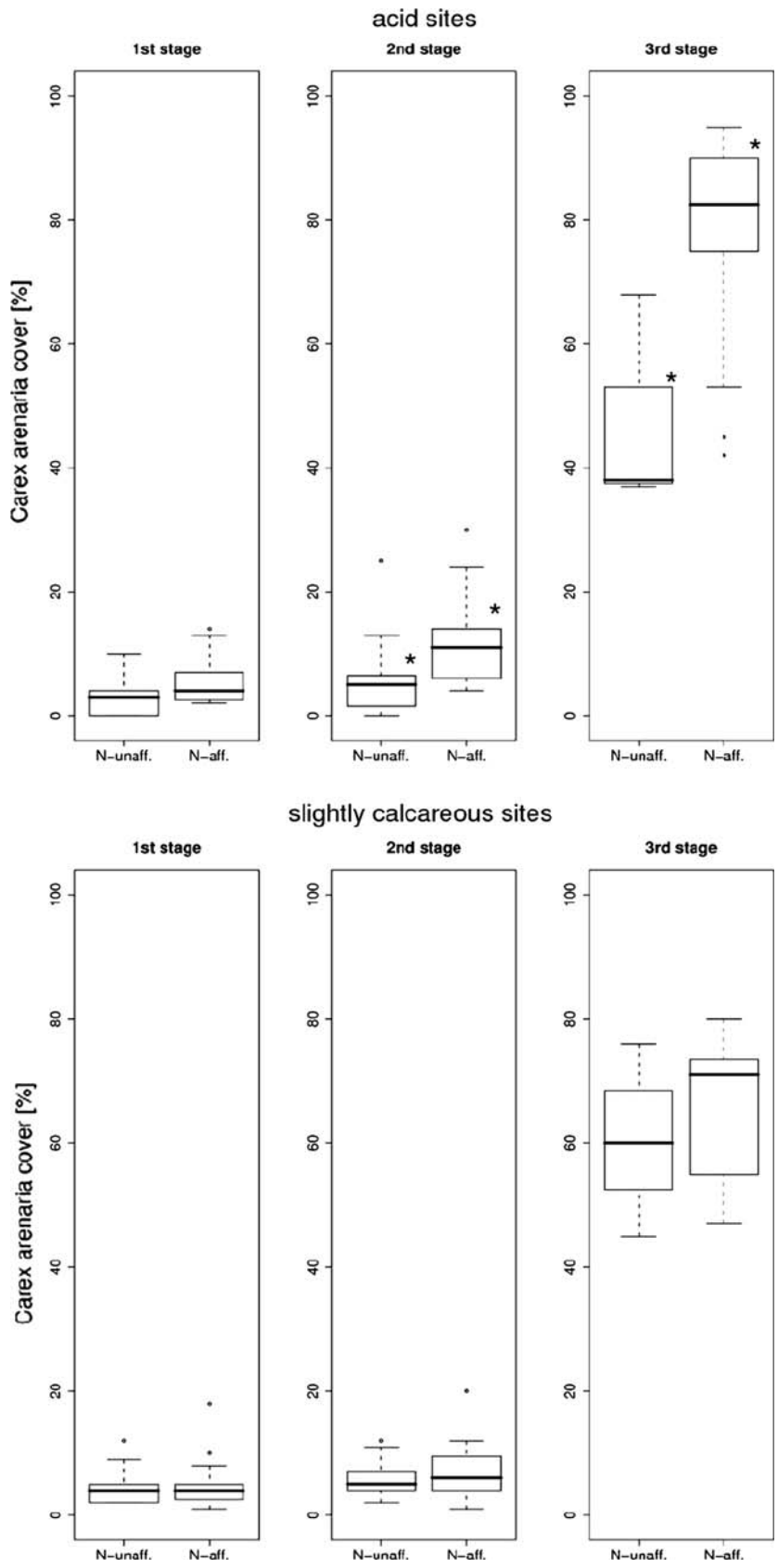


Figure 5. Box and whisker plots for the cover of *Carex arenaria* per plot at acid and slightly calcareous and N-unaffected versus N-affected sites for the three different vegetation succession stages. Significant differences ($P < 0.05$) between N-unaffected and N-affected sites are shown with an asterisk (*).

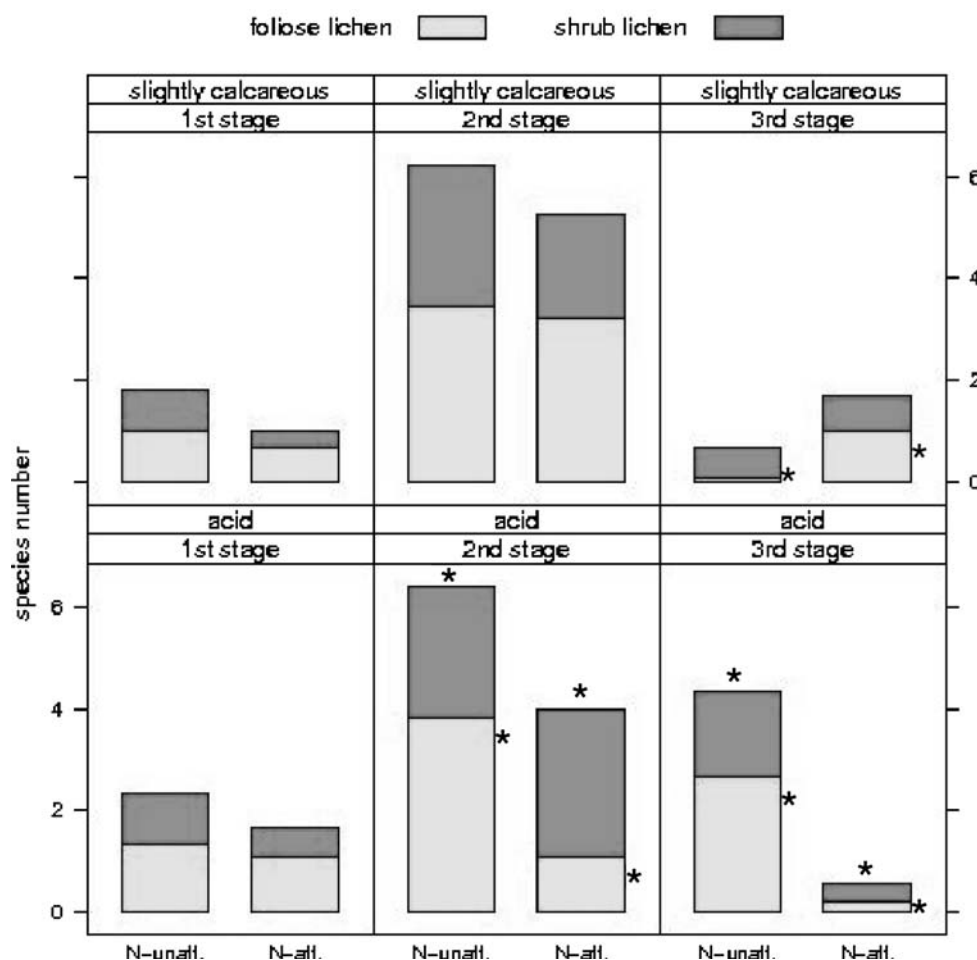


Figure 6. Species number of foliose, shrub and total lichen (per 4 m² plot) at acid and slightly calcareous and N-unaffected versus N-affected sites for the three different vegetation succession stages. Significant differences ($P < 0.05$) between N-unaffected and N-affected sites are shown with an asterisk (*) beside each bar for different lichen groups and on the top of the bar for total lichen species number (foliose and shrub lichen data pooled).

a lower demand for N, and therefore the net availability of N for vascular plants, for example, *Carex*, seems to be higher.

Al/Ca-ratios as well as available Fe concentrations are higher at N-affected, acid sites compared to N-unaffected sites. *Carex* which is more abundant at these sites seems not to be hampered by these high metal concentrations, though metal toxicity might inhibit the growth of many typical plants of dry, lichen-rich dune grasslands. In addition, *Carex* can grow taller and has greater growth plasticity (Tietema 1981) than typical dry dune grassland species, and can thus dominate the system.

A N/P-ratio of 7–12 indicates that *Carex* is far from being P-limited and still N-limited (Koerselman and Meuleman 1996). This is supported by the total tissue concentration of P in *Carex*. P-limitation and critical values depend on plant species and plant tissue part, but a tissue concentration of 50–60 $\mu\text{mol P g}^{-1} \text{ dw}$, which is the range for *Carex* at these 19 sites, is critical but not severely limiting (Troelstra and others 2001).

Carex arenaria is therefore assumed to find optimal growth conditions in Baltic dune sites with high N deposition as it is not hampered by low pH conditions, but is favored by increased availability of its limiting nutrient, nitrogen. With its larger structure, *C. arenaria* can probably comb out more atmospheric N via its leaves as *Ammophila arenaria* does (Heil and others 1988). The N-content and N/P-ratio of *Carex* increase with rising N-deposition mainly at older successional stages, but no correlation was apparent between different soil N-forms and *Carex* tissue N. At older successional stages, atmospheric N may therefore contribute more to the nutrition of *Carex* than soil-derived N. With its extensive rhizome network, *C. arenaria* can exploit and transport temporally and spatially widespread resources nearly all year round (Noble and Marshall 1983; D'Hertenfeld and Falkengren-Grerup 2002; D'Hertenfeld and Jonsdottir 1999) and thereby efficiently use the surplus nitrogen supply. *C. arenaria* finally wins the competition under higher N-loads. Organic matter content above 1.0–1.5% and pH below 4.0 might be the threshold

Table 5. Species Richness for Different Life Forms Separated for Deposition Classes, pH Classes and Succession Stages

	N-unaffected			N-affected		
	First	Second	Third	First	Second	Third
Acid sites						
Foliose lichens	1.3 (0.5)	3.8 (0.8)	2.7 (1.2)	1.1 (0.4)	1.1 (0.3)	0.2 (0.2)
Shrub lichens	1.0 (0.4)	2.6 (0.4)	1.7 (0.9)	0.5 (0.3)	2.9 (0.4)	0.4 (0.2)
All lichens	2.3 (0.9)	6.4 (0.7)	4.3 (1.5)	1.6 (0.5)	4.0 (0.5)	0.6 (0.2)
Mosses	0.8 (0.2)	1.8 (0.3)	1.3 (0.3)	1.0 (0.3)	1.4 (0.2)	0.6 (0.3)
Grasses	2.6 (0.4)	3.0 (0.4)	3.7 (0.3)	2.9 (0.4)	3.4 (0.3)	2.0 (0.2)
Forbs	1.6 (0.6)	3.1 (0.4)	2.3 (0.7)	0.2 (0.1)	3.7 (0.6)	0.9 (0.3)
Total	7.2 (1.2)	14.2 (1.1)	11.7 (2.4)	5.7 (0.9)	12.5 (0.7)	4.1 (0.7)
Slightly calcareous sites						
Foliose lichens	1.0 (0.5)	3.5 (0.8)	0.1 (0.1)	0.7 (0.3)	3.2 (0.7)	1.0 (0.5)
Shrub lichens	0.8 (0.3)	2.8 (0.5)	0.6 (0.2)	0.3 (0.2)	2.1 (0.4)	0.7 (0.3)
All lichens	1.8 (0.7)	6.2 (1.0)	0.7 (0.2)	1.0 (0.4)	5.3 (0.8)	1.7 (0.8)
Mosses	1.0 (0.3)	3.4 (0.4)	1.9 (0.4)	0.9 (0.2)	2.1 (0.2)	1.5 (0.3)
Grasses	2.5 (0.2)	3.7 (0.3)	2.9 (0.4)	3.3 (0.3)	4.1 (0.3)	2.9 (0.2)
Forbs	1.7 (0.4)	6.3 (0.7)	3.5 (0.5)	2.1 (0.4)	5.3 (0.3)	3.3 (0.4)
Total	6.9 (0.8)	19.6 (1.3)	8.9 (1.0)	7.3 (0.7)	16.7 (1.1)	9.4 (1.1)

Values are mean plus 1 SE in brackets; per 4 m² plot.

Significant differences ($P < 0.05$) between deposition classes (N-affected and N-unaffected) are shown in bold. Foliose lichens are lichens with basal thalli mainly flat on the ground like *Cetraria* spp., *Cladonia glauca* or *Cl. foliacea*, *Hypogymnea physodes* and *Peltigera* spp. Shrub lichens are *Cladonia* spp. with no substantial basal thalli and only upwards growing structures, for example, *Cl. arbuscula*, *Cl. furcata*, or *Cl. uncialis*.

conditions for a system shift to a vegetation dominated by *Carex*. Similar shifts in competitive relationships at increasing N-levels have been reported from heathlands and coniferous forests (Aerts and others 1990; Berendse and Aerts 1984; Heil and Bruggink 1987; Keller and Redbo-Torstensson 1995). Another possible reason for grass encroachment is lack of (rabbit) grazing (Veer and Kooijman 1997), but rabbits have not been important grazers according to local site managers and nature conservationists within the 19 dune sites around the Baltic Sea during the last one to two decades, and no other herbivores like elk, red deer, roe deer or hare have been reported to have a major impact on these dry grassland dune systems. Thus, grazing can be excluded as an important factor for grass encroachment by *Carex arenaria* in Baltic dunes.

Character of *Carex*-Dominated Vegetation Units

Dominance by *C. arenaria* can vary considerably. Such vegetation units sometimes are species-rich and open (Harilaid or Pavilosta), but sometimes a species-poor, thick grass sward (Korshage or Gellen). The species-rich, more open *Carex*-dominated vegetation is a community known from Baltic dunes since the 1920s (Steffen 1931; Hueck 1932; Paul 1953). At the Curonian Spit, a slightly cal-

careous and N-unaffected site today, Paul (1953) recorded during the late 1930s an average of 12 species in the optimal phase of a *C. arenaria* community (*Cladonia* spp. excluded). This number is still the same today, and, quite remarkably, species assemblages have not changed during the last seven decades. A decreased number of species (9) at slightly calcareous, N-affected Baltic dunes is comparable to Dutch dunes (Veer and Kooijman 1997).

At higher N loadings, *Carex* dominance is characterized by a dense grass sward. With increasing biomass the light availability decreases, and a dense root network is established (Veer and Kooijman 1997). These types of *Carex* units are probably quite persistent. Once a thick grass sward is set up, it is difficult for other plants and even trees to germinate and establish. Most of the few species that are still occurring in this vegetation unit (three on average) are relics of former, more open stages.

The Effects of N-Deposition on Species Richness

In acid dune systems, even N-loads below 8 kg N ha⁻¹ y⁻¹ can have a negative impact, while slightly calcareous sites are still well enough buffered by the carbonate system (Scheffer and Schachtschabel 1998; Blum 2007), and thus no acidifying impact of

Table 6. Multiple Regression Results of Species Groups with the Factors Lichen-N, Al/Ca-ratio and pH

Species group	First stage		Second stage		Third stage	
	Sig. factor (slope)	R^2 , P-level	Sig. factor (slope)	R^2 , P-level	Sig. factor (slope)	R^2 , P-level
Acid sites						
Foliose lichens	ns		Al/Ca-ratio (-) Interaction with pH Interaction with lichen-N	0.4765*	Lichen N (-)	0.2809 [#]
All lichens	ns		Al/Ca-ratio (-) Interaction with pH Interaction with lichen-N	0.4138 [#]	ns	
Grass	ns		ns		Lichen N (-)	0.5860***
Forbs	Lichen-N (+) pH (+) Interaction of both	0.8907***	ns		ns	
Total	ns		pH (+)	0.1952 [#]	Lichen N (-)	0.3143 [#]
Slightly calcareous sites						
Foliose lichens	ns		ns		Al/Ca-ratio (-) interaction with lichen-N	0.2416 [#]
All lichens	ns		pH (-)	0.3708***	ns	
Moss	ns		lichen-N (-) Al/Ca-ratio (-)	0.3567*	ns	
Grass	ns		pH (-)	0.1784 [#]	ns	
Forbs	Al/Ca-ratio (-)	0.1722 [#]	lichen-N (-) Al/Ca (-) interaction of both	0.5313***	ns	
Total	ns		pH (-)	0.2554*	ns	

Significant factors with algebraic sign of the slope, R^2 and P-level are given (*** $P < 0.0001$, * $P < 0.01$, [#] $P < 0.05$, not significant ns) for the different succession stages at acid and slightly calcareous sites separately.

this moderate atmospheric N-deposition is occurring. As the three factors soil Al/Ca-ratio, N-deposition and soil pH are strongly intercorrelated, their impact cannot be separated totally, but their importance can be ranked. Kleijn and others (2008) could not observe any systematic difference between Al/Ca-ratio at growth sites of common and rare species, but rare species occurred at a restricted pH-range. In this study, Al/Ca-ratio and N-deposition (measured as lichen N), showed the largest impact on dry coastal dunes.

Foliose lichens, which have their thalli directly flat on the ground, disappear along with two to three forb and grass species. *Cladonia glauca* and *Festuca polesica* do not grow at all at N-affected sites. Also in other habitats, coverage and species richness of lichens and mosses decrease at higher atmospheric deposition (Boorman and Fuller 1982; Heil and Diemont 1983; Ketner-Oostra and Sykora 2004; Lee and Caporn 1998; Van Tooren and others 1990). At ambient or artificial total N-deposition of 10 kg N ha⁻¹ y⁻¹ total plant species diversity was reduced in American prairie grasslands (Clark and

Tilman 2008), Swedish deciduous forests (Falken-gren-Grerup and Diekmann 2003) and British grasslands (Stevens and others 2004, 2006). Particular to Baltic dunes, there seems to be a decrease in foliose lichens even at moderate deposition loads of less than 8 kg N ha⁻¹ y⁻¹ wet deposition, whereas shrub-thalli lichens do not change their coverage under these conditions. At N-affected field sites, soil pH is lower and Al and Fe are more available than under pristine conditions, and hence, lichen species not adapted to these high concentrations might take up more metals, which may affect their vitality (Hauk and others 2002, 2007).

Recommendations for Coastal Dune Management

Ecological thresholds and early indicators for ecosystem changes are well used tools in nature conservation and management, though research of their practical application to various ecosystems is still needed (Groffman and others 2006). This study

elucidates the impact of low to medium loads of N deposition on coastal dunes. An early indicator for a dune system change due to N-loads of 5–8 kg N ha⁻¹ y⁻¹ may be an increased drop in soil pH between freshly deposited sand (parent material) and the first successional stage. If the total organic matter content (LOI) in the upper mineral soil horizon of lichen-rich, short grasslands (second stage) is above 1–1.5% and the pH is below 4.0–4.5, the system is about to change. Heavy metals become freely available, for example, Al/Ca-ratios above 1 occur and short grasslands are no longer suitable for slow-growing lichen species. In particular *Cladonia* species growing with their thalli flat on the ground decrease their cover. *Cladonia glauca* and *Cl. macilenta* agg. Hoffm. might be good indicator species of an intact system; higher coverages of *Rumex acetosella* or *Dicranum scoparium* in short grasslands indicate the opposite. Slightly calcareous sites are only marginally affected by these relatively low N-loads as the calcium buffer capacity is not completely depleted yet.

Stands of *C. arenaria* are a natural part of the Baltic coastal ecosystem (Steffen 1931; Hueck 1932; Paul 1953). If the coverage of *C. arenaria* per field site does not exceed 30–40% (Remke and others 2009), these stands can be species rich (12–16 species per 4 m² plot). However, *C. arenaria* stands have become more widespread and more dense. At many sites, a dense, species poor (4–6 species per 4 m² plot) grass sward has established over about 60–70% of the area (Remke and others 2009). Because soil conditions have been irrevocably altered, restoration of dune grasslands on such sites should include turf stripping, removal of soil enriched with organic matter (Ah-layer) and addition of low doses of lime (Dorland and others 2004; Symes and Day 2003).

ACKNOWLEDGEMENTS

This work was funded by the Ministry of Education, Science and Culture, Mecklenburg-Pommern, Germany, and the Bargerveen Foundation, The Netherlands. We are grateful to numerous site managers, researchers and nature conservationists, who helped finding adequate field sites and getting research permits. Thanks also to Henriette Subklew for valuable assistance during field work and to Ankie De Vries-Brock, Jelle Eygenstein and Ulrich Möbius for support in the laboratory. We greatly acknowledge Cathy Jenks for checking the language and two anonymous reviewers and the editor for valuable comments.

OPEN ACCESS

This article is distributed under the terms of the Creative Commons Attribution Noncommercial License which permits any noncommercial use, distribution, and reproduction in any medium, provided the original author(s) and source are credited.

REFERENCES

- Anonymous. 2004. State of environment 2004 report. Vilnius: Ministry of Environment of the Republic of Lithuania. http://www.am.lt/VI/en/VI/article.php3?article_id=93.
- Achermann B, Bobbink R. 2003. Empirical critical loads for nitrogen. Berne: SAEFL.
- Aerts R, Berendse F, de Caluwe H, Schmitz M. 1990. Competition in heathland along an experimental gradient of nutrient availability. *Oikos* 57:310–18.
- Aerts R, Bobbink R. 1999. The impact of atmospheric nitrogen deposition on vegetation processes in terrestrial, non-forest ecosystems. In: Langan SJ, Ed. The impact of nitrogen deposition on natural and semi-natural ecosystems. Dordrecht: Kluwer Academic Publishers. p 85–122.
- Berendse F, Aerts R. 1984. Competition between *Erica tetralix* L. and *Molinia caerulea* (L.) Moench as affected by the availability of nutrients. *Acta Oecol* 5:3–14.
- Biermann RW. 1999. Vegetationsökologische Untersuchungen der *Corynephorus canescens*-Vegetation der südlichen und östlichen Nordseeküste sowie der Kattegatinsel Læsø unter besonderer Berücksichtigung von *Campylopus introflexus*. Mitteilungen der Arbeitsgemeinschaft Geobotanik in Schleswig-Holstein und Hamburg.
- Blum WEH. 2007. Bodenkunde in Stichworten, 6. völlig neu bearbeitete Auflage, Stuttgart: Gebr. Borntraeger.
- Bobbink R, Heil GW, Raessen MB. 1992. Atmospheric deposition and canopy exchange processes in heathland ecosystems. *Environ Pollut* 75(1):29–37.
- Bobbink R, Hornung M, Roelofs J. 1998. The effects of air-borne nitrogen pollutants on species diversity in natural and semi-natural European vegetation. *J Ecol* 86(5):717–38.
- Boorman LA, Fuller RM. 1982. Effects of added nutrients on dune swards grazed by rabbits. *J Ecol* 70:345–55.
- Boorman LA, Van der Maarel E. 1997. Dune grasslands. In: Van der Maarel E, Ed. Dry coastal ecosystems. General aspects 2C. Amsterdam: Elsevier. p 323–44.
- Clark CM, Tilman D. 2008. Loss of plant species after chronic low-level nitrogen deposition to prairie grasslands. *Nature* 451(7179):712–15.
- Crawley MJ. 2005. Statistics: An introduction using R. USA: Wiley.
- Crawley MJ. 2007. The R Book. USA: Wiley.
- De Graaf M, Bobbink R, Verbeek P, Roelofs J. 1997. Aluminium toxicity and tolerance in three heathland species. *Water Air Soil Pollut* 98(3–4):229–39.
- D’Hertenfeld T, Falkengren-Grerup U. 2002. Extensive physiological integration of *Carex arenaria* and *Carex disticha* in relation to potassium and water availability. *New Phytol* 156: 469–77.

- D'Hertenfeld T, Jonsdottir IS. 1999. Extensive physiological integration in intact clonal systems of *Carex arenaria*. *J Ecol* 87:258–64.
- Dorland E, van den Berg L, van de Berg A, Vermeer M, Roelofs J, Bobbink R. 2004. The effects of sod cutting and additional liming on potential net nitrification in heathland soils. *Plant Soil* 265(1–2):267–77.
- Eggertsson-Karlström C. 2004. Väder och Vatten 13. Norrköpping: SMHI.
- Ellenberg H. 1996. Vegetation Mitteleuropas mit den Alpen in ökologischer, dynamischer und historischer Sicht. Stuttgart: Ulmer.
- Falkengren-Grerup U, Diekmann M. 2003. Use of a gradient of N-deposition to calculate effect-related soil and vegetation measures in deciduous forests. *For Ecol Manag* 180(1–3):113–24.
- Feistel R, Nausch G, Wasmund N. 2008. State and evolution of the Baltic Sea, 1952–2005 A detailed 50-year survey of meteorology and climate, physics, chemistry, biology, and marine environment. USA: Wiley.
- Fink A. 2007. Pflanzenernährung und Düngung in Stichworten, 6. völlig neu bearbeitete Auflage. Stuttgart: Gebr. Borntraeger.
- Finnern H. 1994. Ad-Hoc-Arbeitsgruppe Boden der Geologischen Landesämter und der Bundesanstalt für Geowissenschaften und Rohstoffe der Bundesrepublik Deutschland. Bodenkundliche Kartieranleitung. Mit 91 Tabellen. Stuttgart: Schweizerbart.
- Frahm JP, Frey W. 1992. Moosflora. Stuttgart: Ulmer.
- Grasshoff K, Johannsen H. 1977. A new sensitive method for the determination of ammonia in sea water. *Water Res* 2:516.
- Groffman P, Baron J, Blett T, Gold A, Goodman I, Gunderson L, Levinson B, Palmer M, Paerl H, Peterson G, Poff N, Rejeski D, Reynolds J, Turner M, Weathers K, Wiens J. 2006. Ecological thresholds: the key to successful environmental management or an important concept with no practical application? *Ecosystems* 9(1):1–13.
- Hauck M, Hesse V, Runge M. 2002. The significance of stemflow chemistry for epiphytic lichen diversity in a dieback-affected spruce forest on Mt Brocken, northern Germany. *Lichenologist* 34(5):415–27.
- Hauck M, Huneck S, Elix JA, Paul A. 2007. Does secondary chemistry enable lichens to grow on iron-rich substrates? *Flora* 202(6):471–8.
- Hegi G, Conert HJ, Eds. 1998. Illustrierte Flora von Mitteleuropa, 1–4. Berlin: Parey.
- Heil G, Diemont W. 1983. Raised nutrient levels change heathland into grassland. *Vegetatio* 53(2):113–20.
- Heij GJ, Schneider T, Eds. 1991. Acidification research in the Netherlands: final report of the Dutch priority programme on acidification. Amsterdam: Elsevier.
- Heil GW, Bruggink M. 1987. Competition for nutrients between *Calluna vulgaris* (L.) Hull and *Molinia caerulea* (L.) Moench. *Oecologia* 73:105–7.
- Heil GW, Werger MJA, De Mol D, Van Dam D, Heijne B. 1988. Capture of atmospheric ammonia by grassland canopies. *Science* 239:764–5.
- Henriksen A. 1965. An automated method for determining low-level concentrations of phosphate in fresh and saline waters. *Analyst* 90:29–34.
- Houdijk A, Smolders A, Roelofs J. 1993. The effects of atmospheric nitrogen deposition on the soil chemistry of coniferous forests in The Netherlands. *Environ Pollut* 80(1):73–8.
- Hueck K. 1932. Erläuterung zur vegetationskundlichen Karte der Lebanehrung (Ostpommern). (Meßtischblatt Lebasee 171 z. T.). Neudamm, Neumann. Beiträge zur Naturdenkmalpflege.
- Jensen A. 1986. Dry coastal ecosystems of Denmark. In: Van der Maarel E, Ed. Dry coastal ecosystems. 2A. Amsterdam: Elsevier. p 183–96.
- Jones MLM, Wallace HL, Norris D, Brittain SA, Haria S, Jones RE, Rhind PM, Reynolds BR, Emmett BA. 2004. Changes in vegetation and soil characteristics in coastal sand dunes along a gradient of atmospheric nitrogen deposition. *Plant Biol* 6:598–605.
- Keller O, Redbo-Torstensson P. 1995. Effects of elevated nitrogen deposition on the field layer vegetation in coniferous forests. *Ecol Bull* 44:227–37.
- Ketner-Oostra R, Sykora KV. 2004. Decline of lichen diversity in calcium poor coastal dune vegetation since the 1970s, related to grass and moss encroachment. *Phytocoenologica* 35(4):521–49.
- Kingston HM, Haswell SJ. 1997. Microwave enhanced chemistry: fundamentals, sample preparation and applications. Washington, DC, USA: American Chemical Society.
- Kleijn D, Bekker RM, Bobbink R, De Graaf MCC, Roelofs JGM. 2008. In search for key biogeochemical factors affecting plant species persistence in heathland and acidic grasslands: a comparison of common and rare species. *J Appl Ecol* 45(2):680–7.
- Koerselman W, Meuleman A. 1996. The vegetation N:P ratio: a new tool to detect the nature of nutrient limitation. *J Appl Ecol* 33(6):1441–50.
- Kooijman AM. 2004. Environmental problems and restoration measures in coastal dunes in The Netherlands. Chapter 15. Coastal dunes, ecology and conservation. In: Martinez ML, Psuty NP, Eds. Ecological studies, Vol. 171. Berlin: Springer. p 243–58.
- Kooijman AM, Besse M. 2002. The higher availability of N and P in lime-poor than in lime-rich coastal dunes in the Netherlands. *J Ecol* 90(2):394–403.
- Kooijman A, Dopheide J, Sevink J, Takken I, Verstraten J. 1998. Nutrient limitations and their implications on the effects of atmospheric deposition in coastal dunes; lime-poor and lime-rich sites in the Netherlands. *J Ecol* 86(3):511–26.
- Kooijman AM, Kooijman-Schouten MM, Martinez-Hernandez GB. 2008. Alternative strategies to sustain N-fertility in acid and calcareous beech forests: low microbial N-demand versus high biological activity. *Basic Appl Ecol* 9(4):410–21.
- Lautenschlager-Fleury D, Lautenschlager E. 1994. Die Weiden von Mittel- und Nordeuropa: Bestimmungsschlüssel und Artbeschreibungen für die Gattung *Salix* L. Basel: Birkhäuser.
- Lajtha K, Schlesinger W. 1988. The biogeochemistry of phosphorus cycling and phosphorus availability along a desert soil chronosequence. *Ecology* 69(1):24–39.
- Lee J, Caporn S. 1998. Ecological effects of atmospheric reactive nitrogen deposition on semi-natural terrestrial ecosystems. *New Phytol* 139(1):127–34.
- Malloch AJC. 1997. Salt spray as a factor influencing dry coastal vegetation. In: Van der Maarel E, Ed. Dry coastal ecosystems. General aspects. Amsterdam: Elsevier. p 411–18.
- Marschner H. 1995. Mineral nutrition of higher plants. London: Academic Press.
- NEG-TAP. 2001. Transboundary air pollution: acidification, eutrophication and ground-level ozone in the UK, National Expert Group on Transboundary Air Pollution, Edinburgh. <http://www.maposda.net/negtap/finalreport.htm>.

- Niedringhaus R, Haeseler V, Janiesch P, editors. 2008. Die Flora und Fauna der Ostfriesischen Inseln—Artenverzeichnisse und Auswertungen zur Biodiversität. Schriftenreihe Nationalparverwaltung Niedersächsisches Wattenmeer, 11.
- Noble JC, Marshall C. 1983. The population biology of plants with clonal growth II. The nutrient strategy and modular physiology of *Carex arenaria*. *J Ecol* 71:865–77.
- Oberdorfer E. 1994. Pflanzensoziologische Exkursionsflora. Stuttgart: Ulmer.
- Olson DM, Dinerstein E, Wikramanayake ED, Burgess ND, Powell GVN, Underwood EC, D'Amico JA, Itoua I, Strand HE, Morrison JC, Loucks CJ, Allnutt TF, Ricketts TH, Kura Y, Lamoreux JF, Wettengel WW, Hedao P, Kassem KR. 2004. Terrestrial ecoregions of the world: a new map of life on earth. *Bioscience* 51:933–8.
- Paul H. 1953. Morphologie und vegetation der kurischen Nehrung. II. Entwicklung der Pflanzendecke von der Besiedlung des Flugsandes bis zum Wald. *Nova Acta Neopoldina N.F* 16:261–378.
- Power S, Ashmore M, Cousins D. 1998. Impacts and fate of experimentally enhanced nitrogen deposition on a British lowland heath. *Environ Pollut* 102(Suppl 1):27–34.
- Provoost S, Bonte D (eds) (2004). *Levende duinen: een overzicht van debiodiversiteit aan de Vlaamse kust*. Mededelingen van het Instituut voor Natuurbehoud 22. Brussel.
- Raab B, Vedin H, Eds. 1995. *Sveriges Nationalatlas—klimat, sjöar och vattendrag*. Höganäs: Bokförlaget Bra Böcker.
- Ratas U, Nilson E., Eds. 1997. *Small islands of Estonia*. In: *Landscape ecological studies issue 5*. Tallinn: Institute of Ecology. 232 pp.
- R development Core Team. 2008. *R: A language and environment for statistical computing*. R Foundation for Statistical Computing, Vienna, Austria. ISBN 3-900051-07-0. <http://www.R-project.org>.
- Remke E, Brouwer E, Kooijman AM, Blindow I, Esselink H, Roelofs JGM. 2009. Even low to medium nitrogen deposition impacts vegetation of dry, coastal dunes around the Baltic Sea. *Environ Pollut* 157(3):792–800.
- Roelofs JGM. 1986. The effect of airborne sulfur and nitrogen deposition on aquatic and terrestrial heathland vegetation. *Experientia* 42(4):372–7.
- Roelofs J, Kempers A, Houdijk A, Jansen J. 1985. The effect of air-borne ammoniumsulfate on *Pinus nigra* var. *maritima* in The Netherlands. *Plant Soil* 84(1):45–56.
- Rothmaler W, Jäger EJ, Werner K. 2002. *Exkursionsflora von Deutschland 3. Gefäßpflanzen: Atlasband and 4. Gefäßpflanzen: Kritischer Band*, 9. völlig überarbeitete Auflage, Berlin; Heidelberg: Spektrum, Akadem. Verlag.
- Scheffer F, Schachtschabel P. 1998. *Lehrbuch der Bodenkunde*. 14., neu bearb. und erw. Aufl., Stuttgart: Enke.
- Søchting U. 1995. Lichens as monitors of nitrogen deposition. *Cryptogamic Bot* 5:264–9.
- Steffen H. 1931. *Vegetationskunde von Ostpreußen*. Jena: Fischer.
- Stevens CJ, Dise NB, Gowing DJG, Mountford JO. 2006. Loss of forb diversity in relation to nitrogen deposition in the UK: regional trends and potential controls. *Global Change Biol* 12(10):1823–33.
- Stevens C, Dise N, Mountford J, Gowing D. 2004. Impact of nitrogen deposition on the species richness of grasslands. *Science* 303(5665):1876–9.
- Stützer A. 1998. Early stages of podzolisation in young aeolian sediments, western Jutland. *Catena* 32(2):115–29.
- Symes NC, Day J. 2003. *A practical guide to the restoration and management of lowland heathland*. Sandy, UK: RSPB.
- Technicon. 1969. *Industrial method 33–69 W, nitrate + nitrite in water*. Technicon Autoanalyser Methodology. Karrytown-New York: Technicon Corporation. pp 1–2.
- Tietema T. 1981. *Ecophysiology of the sand sedge Carex arenaria L.* Ph.D. thesis, University of Utrecht.
- Troelstra S, Wagenaar R, Smant W, Peters B. 2001. Interpretation of bioassays in the study of interactions between soil organisms and plants: involvement of nutrient factors. *New Phytol* 150(3):697–706.
- van den Berg LJJ, Tomassen HBM, Roelofs JGM, Bobbink R. 2005. Effects of nitrogen enrichment on coastal dune grassland: a mesocosm study. *Environ Pollut* 138:77–85.
- van Dijk HFG, Creemers RC, Rijniers JPLWM, Roelofs JGM. 1989. Impact of artificial ammonium-enriched rainwater on soils and young coniferous trees in a greenhouse. Part I—effects on the soil. *Environ Pollut* 62:317–36.
- van Dijk HFG, De Louw MHJ, Roelofs JGM, Verburgh JJ. 1990. Impact of artificial ammonium-enriched rainwater on soils and young coniferous trees in a greenhouse. Part II—Effects on the trees. *Environ Pollut* 63:41–59.
- Van Tooren BF, Odé B, During HJ, Bobbink R. 1990. Regeneration of species richness in the bryophyte layer of Dutch grasslands. *Lindbergia* 16:153–60.
- Veer M, Kooijman AM. 1997. Effects of grass-encroachment on vegetation and soil in Dutch dry dune grasslands. *Plant Soil* 192(1):119–28.
- Walna B, Polkowska Z, Małek S, Mędrzycka K, Namieśnik J, Siepak J. 2003. Tendencies of change in the chemistry of precipitation at three monitoring station 1996–1999. *Polish J Environ Stud* 12(4):467–72.
- Wirth V. 1995. *Flechtenflora. Bestimmung und ökologische Kennzeichnung der Flechten Südwestdeutschlands und angrenzender Gebiete*. Stuttgart: Ulmer.

URLS FOR CLIMATE DATA

- Greifswald—<http://www.worldclimate.com/cgi-bin/grid.pl?gr=N54E013>.
- Duoedde—<http://www.worldclimate.com/cgi-bin/grid.pl?gr=N55E015>.
- Kleipeda—<http://www.worldclimate.com/cgi-bin/grid.pl?gr=N55E021>.
- Łeba—<http://www.worldclimate.com/cgi-bin/grid.pl?gr=N54E017>.
- Liepaja—<http://www.worldclimate.com/cgi-bin/grid.pl?gr=N56E021>.
- Regions København and Nordsjælland—www.dmi.dk/dmi/index/danmark/vejarkiv.htm.