High Functional Diversity is Related to High Nitrogen Availability in a Deciduous Forest – Evidence from a Functional Trait Approach

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Abstract The current study tested the assumption that floristic and functional diversity patterns are negatively related to soil nitrogen content. We analyzed 20 plots with soil N-contents ranging from 0.63% to 1.06% in a deciduous forest near Munich (Germany). To describe species adaptation strategies to different nitrogen availabilities, we used a plant functional type (PFT) approach. Each identified PFT represents one realized adaptation strategy to the current environment. These were correlated, next to plant species richness and evenness, to soil nitrogen contents. We found that N-efficient species were typical for low soil nitrogen contents, while N-requiring species occur at high N-contents. In contrast to our initial hypotheses, floristic and functional diversity measures (number of PFTs) were positively related.
to nitrogen content in the soil. Every functional group has its own adaptation to the prevailing environmental conditions; in consequence, these functional groups can co-exist but do not out-compete one another. The increased number of functional groups at high N-contents leads to increased species richness. Hence, for explaining diversity patterns we need to consider species groups representing different adaptations to the current environmental conditions. Such co-existing ecological strategies may even overcome the importance of competition in their effect on biodiversity.

**Keywords**  Life history traits · Nitrogen heterogeneity hypothesis · Nitrogen homogeneity hypothesis · Plant functional types (PFT) · Species evenness · Species richness

**Plant nomenclature** Wisskirchen and Haeupler (1998) for vascular plants; Koperski et al. (2000) for bryophytes

**Introduction**

Nitrogen as one of the most important plant nutrients has been the focus of many studies on ecosystem response. In industrialized and densely inhabited countries, nitrogen deposition is one of the most important factors influencing ecosystems because fertilizer use became common: The total atmospheric deposition of reduced nitrogen or nitrogen oxides to terrestrial ecosystems has increased from 17 Tg N/year in 1860 to 64 Tg N/year in the 1990s (Galloway et al. 2004). However, during the last decades, measurements of nitrogen deposition in Central Europe show a slight decrease in nitrogen loads per year (Meessenburg et al. 2009), though the actual levels are still greater than the critical loads given by Bobbink and Roelofs (1995). As a consequence of nitrogen deposition, especially in nitrogen-limited ecosystems, changes have occurred in the vegetation composition (Bobbink et al. 1998; Smart et al. 2003; Bernhardt-Römermann et al. 2006a; Römermann 2006, Römermann et al. 2008). Because nitrogen deposition peaks in areas with high human populations, there is a high risk for nitrogen saturation in Central Europe, particularly in nitrogen-limited ecosystems such as forests (Tamm 1991).

Looking at biodiversity changes in forest ecosystems, the herbaceous layer is of outstanding interest due to its higher species richness in comparison to the tree layer. Furthermore, it responds more easily and quickly to changes in N-deposition compared to trees. Trees respond to increased N-deposition with increased growth rates (e.g., Emmett 1999; Sonnleitner et al. 2001), but with slow changes in species composition (Bernhardt-Römermann et al. 2006b).

In a review Gilliam (2006) concluded that the response of the forest herbaceous layer to increased nitrogen availability presents the following general patterns: i) initial increase in cover of the herb layer as a whole; ii) decrease in species richness due to loss of many N-efficient species; iii) decrease in species evenness due to increasing dominance of few high N-requiring species, and iv) decline in forest biodiversity as a result of decreases in both species richness and evenness. As these predictions are general patterns of ecosystem response, it should be possible to find
these either in time (assuming an increase in N-deposition and therefore N-availability) or looking at spatial gradients. By choosing an existing gradient of nitrogen availability, in the current study we test the abovementioned changes in species diversity in a deciduous forest in Germany (Bernhardt-Römermann et al. 2007).

To learn more about these patterns of vegetation response, a plant functional type approach (PFT) was used. Such an approach is based on the assumption that plants with similar ecologically relevant traits respond to environmental changes in comparable ways (McIntyre et al. 1995). A trait is called functional when it impacts plant fitness indirectly via its effects on growth, reproduction, and survival. If the trait shows responses to changes in environmental conditions it is called a response trait; the values or modality taken by a trait at a point of an environmental gradient is called an attribute (Viéville et al. 2007). A group of functional traits is called a plant functional type (PFT), but to be ecological meaningful each PFT should contain the most parsimonious number of traits only (as less traits as possible, but as much as needed; Keith et al. 2007; Bernhardt-Römermann et al. 2008). Using such PFTs we can estimate which functional trait composition is the result of differences in nitrogen availability on the estimated nitrogen gradient. Hence, the applied PFT approach can be used to learn more about patterns of biodiversity: different PFTs represent different adaptations of the co-occurring plant species to the current environmental conditions (complementarity of resource use; Díaz and Cabido 2001). Thus, we may explain high species richness by a high number of PFTs. Following Gilliam (2006) a higher number of PFTs should occur at nitrogen-poor situations in comparison to N-rich situations.

Material and Methods

Study Area

The study area, called Echinger Lohe, is located northeast of Munich (11°43’ E, 47°55’ N, 465 m a.s.l.). It is a small deciduous forest of approximately 24 ha, which is surrounded by intensively used farmland. The mean annual temperature is approximately 9°C and the average annual rainfall about 800 mm. The Echinger Lohe is situated on the “Münchener Schotterebene”, a plain that was formed by homogeneously deposited gravel at the end of the last ice age. Due to intensive agriculture in the vicinity of the forest, the background level of nitrogen pollution is about 15–20 kg/ha (Bayerische Landesanstalt für Wald und Forstwirtschaft 2005). The vegetation type can be characterized as a Quercus petraea-Carpinus betulus forest with Acer pseudoplatanus on pararendzina soil.

Vegetation Sampling

To characterize the actual vegetation composition of the Echinger Lohe, 20 randomly distributed relevés of 15 m × 15 m each were carried out in 2003 (the distance of each relevé is at least 40 m to footpaths or forest edges). This relevé size is recommended by Westhoff and van der Maarel (1973) for sampling the forest floor vegetation.
Covers of vascular plants and bryophytes were estimated using the Braun-Blanquet method (1964) with classes following Pfadenhauer et al. (1986; +: <1%; 1a: 1%–3%; 1b: 3%–5%; 2a: 5%–15%; 2b: 15%–25%; 3: 25%–50%; 4: 50%–75%; 5: 75%–100%) and divided into herb, shrub, and tree layer. Additionally, the overall cover of the herb layer was estimated in the field (using a percentage scale divided into 5% interval classes). For data analyses, the cover of each species was transformed into the mean value of the limits in the corresponding class.

Estimation of the Nitrogen Gradient

Three soil samples were taken from the Ah-horizon of each relevé and pooled for further analyses to characterize the herb layer’s main rooting space. These samples were analyzed following the German standard method for analyses of forest soils (Bodenzustandserhebung im Wald- BZE; Bundesministerium für Ernährung Landwirtschaft und Forsten 1994) for N, C, pH(H2O), NH4+, NO3-, Cl-, H2PO4-, SO4²-, Mg²+, Ca²+, K⁺, Na⁺, Al³⁺, Fe²⁺, and Mn²+. Additionally, the thickness of the Ah-horizon was reported.

To characterize the three vegetation types defined by a cluster analysis, dry nitrogen deposition (NH3 and NO2) was measured using passive samplers (one sampler per vegetation type in 2004). Wet deposition was calculated from the bulk precipitation in an adjacent non-wooded area. The potential nitrogen deposition was calculated by adding values of dry and wet deposition (see Bernhardt-Römermann et al. 2006a).

Estimation of Differences in Species Composition

The vegetation data from the 20 relevés were clustered into groups based on differences in their floristic composition. Relative Euclidean Distance and Ward’s method were applied on transformed vegetation data following the formula \( y = x^{0.25} \) (Wildi 1986), to reduce the weight of dominant species. Each of these vegetation types was characterized by its typical species set by the use of an indicator species analysis (ISA; Dufrene and Legendre 1997). An indicator species analysis assigns indicator values for each species, based on the degree to which they discriminate among groups. The indicator values (IV) range from zero (no indication) to 100 (perfect indication). For each indicator value a randomization test is used to evaluate its statistical significance. The null hypothesis is that the observed IV is no larger than would be expected by chance (i.e., that the species has no indicator value). All these statistics were done using the program PcOrd 5.21 (McCune and Mefford 1999).

Diversity indices (species richness and species evenness) and herb-cover were analyzed. These parameters were calculated including all herbaceous and bryophyte species. We calculated Pearson’s product-moment correlations between the above-mentioned vegetation parameters and the soil nitrogen content and tested for differences between vegetation groups using R 2.9.0 (R Development Core Team 2009) randomization tests. 1,000 randomized samples were taken and the difference to the observed (measured) mean was calculated; if random samples show in less than 0.01% of all permutations a mean lower than the observed, a significant difference was assumed.

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Plant Functional Types

Using a plant functional type approach (PFT), the spatial nitrogen gradient was analyzed to identify nitrogen-induced vegetation differentiation. The analysis combined vegetation, trait, and environmental data and followed the numerical procedure by Pillar (Pillar 1999; Pillar and Sosinski 2003). Data were arranged in three different tables: i) relevés by species matrix, containing species percentage covers, ii) species by traits matrix, and iii) relevés by environmental variables matrix.

Bryophytes were excluded from further analyses because trait data were not available. To characterize the environment, we used those parameters that described the differences in vegetation composition best: the N-content of the soil and the thickness of the Ah-horizon (see below). The species descriptions by traits were taken from different trait databases. The traits cover different dimensions of plant life-history, including persistence and regeneration traits and are summarized in Table 1. All PFT analyses were carried out with the program SYNCSA 2.2.4 (Pillar 2004) using the polythetic clustering option. This program searches for the most relevant traits as part of the analyses themselves, therefore no further justification if the trait is relevant or not, is needed. In the following only a brief overview of the method is given, for details see Pillar and Sosinski (2003).

The iterative algorithm was used to evaluate all possible groups of species by cluster analyses with different subsets of traits. For each of these possible solutions the congruence between variation in vegetation composition and in the environment was measured using matrix correlation. The latter was calculated between the vegetation composition distances described by the species groups (using Chord’s distance) and the site’s distances described by the environment (using Euclidean distance after standardization of the variables by unit length). A trait subset was identified from all possible solutions for which the correlation (here called congruence) between the matrices was maximal.

In the second step, the PFTs defined by this optimal trait subset were used to describe the relevés, resulting in a PFT by relevés matrix. These results were visualized using a principal coordinates analysis (PCoA, Gower 1966, with Chord’s distance).

For each plant functional type, the ecological performance was calculated by summing up the coverage of all species belonging to the respective PFT per relevé. Subsequently, the mean coverage per PFT was taken over all relevés. This value was relative with respect to the total cover of all species.

Results

As pointed out in Bernhardt-Römermann et al. (2007) the most important soil parameters for explaining the variance between the different vegetation types were the soil nitrogen content and the thickness of the main rooting horizon. The vegetation of the Echinger Lohe can be separated into three vegetation types (Bernhardt-Römermann et al. 2007). Table 2 shows the results of the indicator species analysis. It revealed that the following species characterize the different vegetation types: vegetation type 1 included the low-nitrogen tolerant (see Ellenberg N-values in Table 2) species Carex montana, Rhytidiadephus triquetrus, Fragaria
vesca, Rubus saxatilis, Convallaria majalis, and Vincetoxicum hirundinaria. Type 3, however, was characterized by the occurrence of species that are typical for high N-availability (see Ellenberg N-values in Table 2): Corydalis cava, Asarum europaeum, Heracleum spongylion, Arum maculatum, Allium carinatum, Elymus caninus, Stachys sylvatica, Gagea lutea, Aegopodium podagraria, Chaerophyllum aureum, Eurchynichium swartzii, Polygonum multiflorum, and Plagiommium undulatum. For vegetation type 2 only Paris quadrifolia was found as a significant indicator species. These patterns and interpretations of the vegetation differentiation were consistent with the results of the soil analyses. As stressed by Bernhardt-Römermann et al. (2007) the actual gradient in vegetation differentiation was closely

| Trait | Abr. | Type | Trait states | State classes according to | Data origin |
|-------|------|------|--------------|----------------------------|-------------|
| Plant height (max.) | MH | 2 | 1: <30 cm; 2: 30–60 cm; 3: >60 cm | Kleyer (1995) Rothmaler (2002) |
| Rosettes | RO | 1 | 1: erosulate plant; 2: hemirosette plant; 3: rosette plant | Klotz and Kühn (2002) Klotz et al. (2002) |
| Leaf anatomy | AN | 1 | 1: scleromorphic; 2: mesomorphic; 3: hygromorphic | Klotz and Kühn (2002) Kleyer (1995); Klotz et al. (2002) |
| Leaf persistence | LP | 1 | 1: persistent green; 2: spring green; 3: summer green | Ellenberg et al. (2001) Klotz et al. (2002); BfN (2003) |
| Leaf form | LF | 1 | 1: grass-like; 2: long leaf; 3: simple; 4: full; 5: lobate; 6: pinnatifid; 7: digitate; 8: palmate; 9: pinnate; 10: bipinnate | Klotz and Kühn (2002) Oberdorfer (2001); Klotz et al. (2002) |
| Vegetative spread | VS | 1 | 1: <100 mm, CLOPLA1 (numbers 1, 2, 4, 6, 7, 9, 12, 13, 15–19); 2: ≥100 mm, CLOPLA1 (numbers 3, 5, 8, 10, 11, 14) | Kahmen (2004) Klimeš et al. (1997) |
| Plant persistence | PP | 1 | 1: annual; 2: perennial, not clonal, CLOPLA1 (numbers 1, 2, 4, 12, 16–19); 3: perennial, clonal, CLOPLA1 (numbers 3, 5, 11, 13–15) | Kahmen (2004) Klimeš et al. (1997) |
| Seed mass | MA | 2 | 1: ≤0.2; 2: 0.21–0.5; 3: 0.51–1; 4: 1.01–2; 5: 2.01–10; 6: >10 (mg/1,000 seeds) | Grime et al. (1988) Flynn et al. (2004) |
| Pollination | PO | 1 | 1: selfing; 2: by insects; 3: by wind | Klotz et al. (2002); BfN (2003) |
| Start of flowering | SF | 1 | 1: before or in May; 2: after May | Kahmen (2004) Rothmaler (2002); BfN (2003) |
| Duration of flowering | DF | 1 | 1: 1–2 months; 2: ≥3 months | Kahmen (2004) Rothmaler (2002); BfN (2003) |
related to differences in soil nitrogen content (mean N content in %±s.e.; type 1: 0.76±0.04; type 2: 0.72±0.04; type 3: 0.94±0.03) and to thickness of the main rooting horizon (mean Ah-horizon thickness in cm±s.e.; type 1: 6.7±0.5; type 2: 7.5±0.4; type 3: 11.6±0.7); all other soil parameters were found to be irrelevant. Looking at the annual N-deposition, again an increase from vegetation type 1 to 3 was found (type 1: 17.6 kg/ha, type 2: 19.9 kg/ha; type 3: 21.5 kg/ha).

The results of the correlation analyses between herb layer cover, evenness, species richness and soil N-content showed significant positive relationships for all three correlations (Table 3). Between the three vegetation types we found significant differences in herb layer cover, evenness, and species richness (Table 3). The herb layer cover continuously increased from vegetation type 1 to 3. For evenness no differences between type 1 and 2 were found, whereas vegetation type 3 showed significantly higher values. Richness peaked in vegetation type 3, but only in vegetation type 2 was it significantly lower.

**Plant Functional Types**

All solutions of possible trait combinations that characterize the different plant types included traits that were related to biomass development, plant persistence (for

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**Table 2** Results of the indicator species analysis (ISA). Only non-tree species having significant indicator values are shown. For each species, its indicator value (IV) and typical vegetation type is reported. Additionally, Ellenberg-values for nitrogen (N) are given.

| Species                          | Type | IV  | P-value | N   |
|----------------------------------|------|-----|---------|-----|
| Carex montana                    | 1    | 98.3| 0.000   | 3   |
| Rhytidiadepus triquetrus         | 1    | 87.8| 0.005   |     |
| Fragaria vesca                   | 1    | 67.5| 0.010   | 6   |
| Rubus saxatilis                  | 1    | 58.3| 0.030   | 4   |
| Convallaria majalis              | 1    | 79.2| 0.037   | 4   |
| Vincetoxicum hirundinaria        | 1    | 50.0| 0.044   | 3   |
| Paris quadrifolia                | 2    | 65.5| 0.033   | 7   |
| Asarum europaeum                 | 3    | 87.5| 0.000   | 6   |
| Heracleum sphondylium           | 3    | 87.5| 0.000   | 8   |
| Arum maculatum                   | 3    | 94.8| 0.001   | 8   |
| Corydalis cava                   | 3    | 87.5| 0.002   | 8   |
| Allium carinatum                 | 3    | 68.4| 0.004   | 2   |
| Elymus caninus                   | 3    | 75.0| 0.004   | 8   |
| Stachys sylvatica                | 3    | 67.8| 0.011   | 7   |
| Gagea lutea                      | 3    | 64.3| 0.013   | 7   |
| Aegopodium podagraria            | 3    | 61.3| 0.028   | 8   |
| Chaerophyllum Aureum             | 3    | 50.0| 0.042   | 9   |
| Eurhynchium schwartzii           | 3    | 50.0| 0.043   |     |
| Polygonum multiflorum            | 3    | 58.8| 0.043   | 5   |
| Plagiomnium undulatum            | 3    | 50.0| 0.046   |     |
example plant height, leaf form or leaf persistence) or flowering traits (start or
duration of flowering); see Table 4 for an overview of different solutions and
included traits. Even though congruence differs only slightly between the first five
solutions (Table 4), we included the second solution proposed by the algorithm in
further analyses, because it represented the most parsimonious solution with respect
to numbers of PFTs needed for describing the vegetation differentiation (n=7). This
combination contained the traits leaf persistence, plant height, start of flowering, and
seed mass.

Figure 1 showed the results of the PCoA of the relevés described by the extracted
PFTs. The relevés were arranged from lower to higher nitrogen availability, where
axis one described the nitrogen availability (correlation with soil N-content: r²=0.82,
P<0.001). The performances of PFTs number 2, 3, 5, and 6 were associated with
that gradient. From these, PFTs number 2, 3, and 5 formed a response group to high
nitrogen availability, whereas PFT 6 characterized lower nitrogen contents (Fig. 1
and Table 5). The other PFTs (1, 4 and 7) show no response to that N-gradient and
were omitted from further analyses.

The trait states of these four responding PFTs were indicated in Table 5. The
plants of PFTs 2 and 5 were tall and flower late in the year. Those of PFT 2 were
permanent green with seed masses between 1 and 2 mg, whereas PFT 5 contained
summer green plants with lighter seeds (0.5 to 1 mg). Differently, the plants of PFT

Table 3  Diversity indices and cover of the herb layer for the three vegetation types (1 – low; 2 – intermediate;
3 – high nitrogen availability). Additionally, correlations with N-content of the soil are presented. Means and
standard errors are given. Different letters indicate significant differences within each category

|                | Correlation with soil N-content (d.f.=16) | Vegetation type 1 (n=8) | Vegetation type 2 (n=4) | Vegetation type 3 (n=8) |
|----------------|------------------------------------------|------------------------|------------------------|------------------------|
|                | t-value | P-value | r-value | Mean±s.e. | Mean±s.e. | Mean±s.e. |
| Richness       | 2.50    | 0.022   | 0.51    | 30.9±2.36 (ab) | 24.5±2.84 (a) | 33.3±1.39 (b) |
| Evenness       | 2.27    | 0.036   | 0.47    | 0.49±0.03 (a) | 0.48±0.06 (a) | 0.63±0.03 (b) |
| Cover          | 2.52    | 0.021   | 0.51    | 65.0±3.41 (a) | 66.3±6.25 (ab) | 77.5±2.67 (b) |

Table 4  Trait subsets with highest values of congruence between the vegetation composition described by
plant functional types with corresponding plant traits and the relevé sites described by the environmental
variables, soil N-content and thickness of the Ah-horizon. Only optimal numbers of PFTs are presented.
See Table 2 for explanation of trait abbreviations. The trait set used for further analyses is indicated by
bold letters

| Number | Congruence | No. of PFTs | Subset of traits |
|--------|------------|-------------|------------------|
| 1      | 0.701      | 12          | MH MA PP DF      |
| 2      | 0.684      | 7           | LP MH MA SF      |
| 3      | 0.675      | 12          | LP AN MH MA SF DF LF |
| 4      | 0.671      | 11          | LP MH MA SF DF   |
| 5      | 0.671      | 11          | LP MH SF DF      |
Table 5 Trait states of the seven PFTs found by the matrix correlation. For the analysis the traits leaf persistence, height of plant, start of flowering, and seed mass were used. Those PFTs found best for describing the environmental gradient using the PCoA (Fig. 1) were marked in bold letters. PFTs showing a positive correlation with the first axis of the PCoA were grouped together in the response group 1 (high nitrogen availability), and those with a negative correlation were put into response group 0 (low nitrogen availability).

| PFT  | Response group | Leaf persistence | Height of plant | Seed mass | Start of flowering |
|------|----------------|-----------------|-----------------|-----------|-------------------|
| 1    |                | persistent green| $\leq 30$ cm    | 1–2 mg    | before or in May  |
| 2    | 1              | persistent green| $\gt 60$ cm     | 1–2 mg    | after May         |
| 3    | 1              | spring green    | $\leq 30$ cm    | 2–10 mg   | before or in May  |
| 4    |                | summer green    | 30–60 cm        | 2–10 mg   | after May         |
| 5    | 1              | summer green    | $\gt 60$ cm     | 0.5–1 mg  | after May         |
| 6    | 0              | summer green    | 30–60 cm        | 2–10 mg   | before or in May  |
| 7    |                | persistent green| $\gt 60$ cm     | 1–2 mg    | before or in May  |

3 were small, spring green, early-flowering, and had heavy seeds. PFT 6, the only representative of the response group 0 (i.e., indicating low nitrogen availability), comprised summer green, medium-sized species that flower early in the year and had heavy seeds. In total, more than 74% of all herbaceous species occurring in the Echinger Lohe were described using one of these PFTs (Table 6).

Discussion

The results of the current study clearly showed that the vegetation differentiation in the Echinger Lohe is closely related to a gradient in nitrogen availability. Next to soil N-content we found differences in the thickness of the Ah-horizon that are related to differences in plant available water. However, it is well known from literature (e.g., Ellenberg 1996) that additional water or nutrient supply has a similar effect on plant growth, as both induce advanced nutrient uptake (if there is a sufficient minimum of water available). Thus, the influence of the two factors, soil N-content and increased
water availability, cannot be separated, but both are clearly related to nutrient uptake (Bernhardt-Römermann et al. 2007).

As already outlined in the introduction, vegetation development under increased nitrogen deposition and, therefore, availability can be divided into several steps: First, an increase in total vegetation cover, followed by a decrease of the nitrogen-efficient species (which are typical for conditions with low nitrogen availability) can be expected (Gilliam 2006). When describing the development of the ecosystems under increased nitrogen deposition, Gilliam (2006) assumed high species richness in the original stage, including species on N-poor and N-rich soils. These species coexist due to the spatial heterogeneity in nitrogen availability (nitrogen heterogeneity hypothesis: Ricklefs 1977; Tilman 1984; Rosenzweig and Abramsky 1993; Tilman and Pacala 1993). The increase in total vegetation cover is linked to the removal of the growth-limiting factor nitrogen, i.e., additional nitrogen is added to the system. Decreasing numbers of nitrogen efficient species can be explained by the increase of N-requiring species that out-compete N-efficient species (Diekmann and Falkengren-Grerup 2002). The results of the current study are consistent with the described development because we could show a close positive relationship between vegetation cover and N-content of the soil. Furthermore, vegetation type 1, which is typical for nitrogen-poor situations, is characterized by several N-efficient species (here indicated by Ellenberg N-indicator values).

Subsequently, Gilliam (2006) suggested a decline in evenness due to the dominance of relatively few N-requiring species. Together with the decrease in N-efficient species, this leads finally to reduced biodiversity (in richness and evenness). Our results show a clear differentiation in species typical for nitrogen-rich and poor environments; under poor conditions N-efficient species dominated, in soils with a high nitrogen content N-requiring species dominated. Nevertheless, we did not find a decrease in total species richness and evenness with increasing N-content; rather we found a positive relationship between N-content and species richness as well as evenness. We can explain this discrepancy to the aforementioned predictions of

Table 6  Species belonging to the different plant functional types (PFT). Numbers of PFTs and response groups correspond to those given in Table 5. Additionally, the relative ecological performance (REP, in % of mean total vegetation cover) is given for each PFT

| PFT | Response group | Species                                                                 | REP   |
|-----|----------------|-------------------------------------------------------------------------|-------|
| 2   | 1              | *Allium carinatum*, *Lamium montanum*, *Geum urbanum*                   | 11.1  |
| 3   | 1              | *Anemone nemorosa*, *Arum maculatum*, *Corydalis cava*, *Gagea lutea*, *Ranunculus ficaria* | 8.2   |
| 5   | 1              | *Brachypodium sylvaticum*, *Campanula trachelium*, *Aegopodium podagraria*, *Allium scorodoprasum* subsp. *rotundum*, *Lilium martagon*, *Stachys sylvatica*, *Poa nemoralis*, *Chaerophyllum aureum*, *Elymus caninus*, *Heracleum sphondylium* | 17.7  |
| 6   | 0              | *Carex montana*, *Convallaria majalis*, *Lamium maculatum*, *Maianthemum bifolium*, *Melica nutans*, *Polygonatum multiflorum*, *Paris quadrifolia*, *Viola hirta*, *Vincetoxicum hirundinaria*, *Primula veris*, *Viola mirabilis*, *Mercurialis perennis*, *Ranunculus lanuginosus* | 26.7  |
Gilliam (2006) using a functional trait approach, as life-history traits of species are strongly related to the processes of vegetation differentiation (Diekmann and Falkengren-Grerup 2002; Lavorel and Garnier 2002).

Assuming that the assemblage of species in most forest ecosystems is mainly determined by competition and stress tolerance (Grime 1979), ecosystems contain different plant strategies to compete for ecologically relevant resources. Different functional traits can be seen as strategies of resource use in which species of different PFTs show a higher degree of complementarity than those of the same PFT (Díaz and Cabido 2001). The more different PFTs co-occur in an ecosystem, the more ecologically relevant strategies co-exist and the higher the diversity may be (Mason et al. 2005). Nevertheless, most species are not equally distributed in their dominance patterns in ecosystems (Wilson and Agnew 1992; Grime 1998). The meaning of several realized strategies becomes even more relevant if these are dominated by only a few species: Evenness largely depends on the number of plant functional types under certain environmental conditions, i.e., in this case on nitrogen stages (Díaz and Cabido 1997; Hector et al. 1999; Chapin et al. 2000; Loreau et al. 2001). In the current study the response group 1, characterizing vegetation of high nitrogen availability, is composed of three PFTs. In comparison, response group 0, which is typical for nitrogen-poor conditions, consists of only one PFT. Within response group 1, it was possible to identify three main strategies: The plants of the first two strategies are described by two PFTs (PFT 2 and 5); both represent tall, persistent or summer-green species that flower during the summer. In contrast, plants of the second strategy (PFT 3) are small species that start flowering early in spring. As also shown by Diekmann and Falkengren-Grerup (2002), tall species in the herb layer benefit from high amounts of nitrogen; for these species, the decreased light conditions (when the tree layer is foliated) can be compensated by better nutrient supply (Peace and Grubb 1982; Peace 1984). The small species belonging to PFT 3 can also be found in nutrient-rich forests (Al-Mufti et al. 1977; Rogers 1982; Host and Pregitzer 1991; Rothstein and Zak 2001), showing high rates of photosynthesis depending on high water and nutrient availability (Grime 1979). These species, however, evade competition by finishing their life cycle before the start of the foliation of the trees.

Within response group 0 (stands with lower levels of plant-available nitrogen) only one single PFT could be identified, which is characterized by medium-sized and summer-green plant species. Because of the lower thickness of the soil layer, dryness during summer is common (personal observation) and nitrogen availability is limited, which causes low cover and height of the vegetation. Stress is the most relevant factor for this vegetation type, whereas competition for growing space or light is less important; the forests are dominated by summer-green or persistent species (Grime 1994).

The PFT-approach used in the current study allows us to get information on how plant species use different strategies to compete for environmental resources. High species richness and species evenness under nitrogen-rich conditions can be clearly attributed to the higher number of functional adaptations to the current environment: As the number of PFTs increases, the species number increases, but the relative performance of each PFT decreases (the relative performance of each of the three PFTs under nutrient-rich conditions: <18% vs relative performance of the PFT 6...
under nutrient-poor conditions: 26.7%; see Table 6). Taking into account that each PFT is dominated by a few species (i.e., has a relatively low evenness) it becomes clear that the co-existence of several PFTs involves at the same time a co-existence of several “dominant” plant species: The higher biodiversity (richness as well as evenness) under nitrogen-rich conditions is strongly related to a higher variety of strategy types.

Thus, our results clearly showed contrasting results with respect to Gilliam’s (2006) hypotheses: he hypothesized that chronically elevated nitrogen deposition leads to a decline in biodiversity due to a decrease of the spatial heterogeneity of nitrogen (nitrogen homogeneity hypothesis); in other words: nitrogen is made uniformly available in contrast to patchiness in more oligotrophic situations. Under these conditions of increased N-deposition, vegetation changes are due to a switch from a coexistence of strong and weak competitors to a dominance of highly competitive species only. However, in the current study we found at high nitrogen availability an increased number of strategies of adaptation to the current environmental conditions – co-existence of species is possible due to a differentiation in use of ecological relevant recourses. Such co-existing ecological strategies may overcome the importance of competition in their effect on biodiversity.

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