INTRODUCTION

In the face of unprecedented loss in global diversity, the relationship between diversity and stability has again become a major focus of research in ecology. Whereas evidence accumulates for consistent diversity effects on the magnitude of processes (see e.g. Cardinale et al. 2006, Hillebrand & Matthiessen 2009), there is ongoing debate as to whether diversity induces certain aspects of stability.

Two aspects of stability have been to date analysed separately, although they are not fully distinct from each other: variability over time and response to disturbance. In the context of the diversity–stability hypothesis (DSH), diversity is suggested to reduce the temporal fluctuation in aggregate ecosystem processes by compensatory dynamics (Tilman 1996, Cot-
tingham et al. 2001), statistical averaging (Doak et al. 1998), or by decreasing the synchrony of population dynamics (Roscher et al. 2011). This aspect of the DSH is comparably well underpinned by theory, e.g. by the insurance model (Yachi & Loreau 1999), which bases diversity effects on the fact that diverse communities are more likely to show compensatory dynamics than communities of low biodiversity. Although compensatory dynamics in response to environmental fluctuation has been analysed in natural systems (Vasseur & Gaedke 2007, Winfree & Kremen 2009) and experiments (Gonzalez & Descamps-Julien 2004, Downing 2008), there is considerable debate as to whether compensatory dynamics generally prevail in ecosystems (Houlahan et al. 2007, Valone & Barber 2008).

Regarding the second aspect, the response to disturbance, it has been suggested that the importance of biodiversity for community response to a pulse perturbation depends on the resistance (the ability to withstand disturbance) and resilience (the ability to recover from disturbance) of the species within the community (Pfisterer & Schmid 2002, van Ruijven & Berendse 2010). In this context, disturbance is defined as a temporally explicit phenomenon that reduces community biomass by causing its partial or total destruction (sensu Grime 1979). The prediction of how biodiversity affects these 2 aspects of stability is not straightforward as each potentially requires different species traits. These traits enable the species to resist or tolerate the disturbance or to grow rapidly after the disturbance. Trade-offs between growth and defendedness can be expected (Fine et al. 2006), as adaptations to resist disturbances potentially come at a cost to growth rate, which defines whether a species can contribute quickly to recovery. The relative proportion of sensitive, tolerant and resistant species may, therefore, foster different diversity–stability relationships in the response to disturbance. Consequently, experimental studies (e.g. Pfisterer & Schmid 2002, Moorthi et al. 2008) remained inconclusive with respect to the presence of general biodiversity effects on stability following pulse perturbations.

The overall relationship between biodiversity and stability after disturbance will thus depend on how the traits that relate to resistance and recovery are distributed within the community. This suggests that the dominance structure (evenness) of communities may be as important for community stability as richness (Hillebrand et al. 2008, Sasaki & Lauenroth 2011). We propose that population dynamics of dominant species (low evenness) determine the response of the entire community, if these species display high resistance or resilience. If dominant species are highly sensitive and recover slowly, compensatory growth of rare species is needed to maintain or recover high levels of ecosystem performance. In this case, higher evenness increases stability. Thus, the relationship between diversity and response to disturbance is largely determined by interdependencies between the number of species present and their identity and degree of dominance.

In addition to the unclear separation between different aspects of stability and diversity, studies within the framework of the DSH have focussed entirely on the effects of single disturbances, although natural systems are commonly affected by multiple aspects of disturbance (Liiri et al. 2002). As stressors (= disturbances) are expected to potentially interact with non-additive consequences for biomass reduction, the stability of communities that are subjected to multiple stressors cannot be extrapolated from studies investigating single stressors (Breitburg et al. 1998). Vinebrooke et al. (2004) proposed 3 scenarios of multiple stressor effects in the framework of the DSH: species’ tolerances to multiple stressors may be randomly distributed, or they may be positively or negatively correlated. A positive correlation would result in higher stability, because species that survive the impact of Stressor A are likely to tolerate Stressor B (stressor-induced community tolerance). Conversely, a negative correlation between species’ tolerances to multiple stressors would lead to a strong decrease in stability (stressor-induced community sensitivity). If species’ tolerances to stressors are randomly distributed and unrelated, exposure to Stressor A is not likely to influence the effect of Stressor B.

The objective of our study was to investigate the effects of species richness and species identity on the stability of communities that are affected by multiple stressors. We performed 2 experiments using various strains of freshwater phytoplankton affected by combinations of 2 artificial stressors: pH reduction and grazing. First, we analysed the response of these strains to both stressors, as well as their growth rates, in a monoculture experiment. In the subsequent diversity experiment, we simulated 2 different scenarios of community composition using a combinatorial design. The first scenario (Combination A) focussed on the hypothesis that stability depends solely on the population dynamics of a dominant species with functional traits allowing its persistence after disturbance. This species was present at all levels of species richness (2, 4 and 8 species). The
second scenario (Combination B) tested the hypothesis that, in the absence of such a dominant and highly functional species, the richness of species determines the response of a community to multiple stressors.

MATERIALS AND METHODS

Experimental conditions

Eleven phytoplankton strains represented the species pool of our study: Cylindrospermum sp. (Cyl), Cyclotella sp. (Cyc), Melosira sp. (Mel), Fragilaria capucina (Fra), Cryptomonas pyrenoidifera (Cry), Tetraselmis sp. (Tes), Chlamydomonas tericola (Chl), Pediasastrum sp. (Ped), Oocystis sp. (Ooc), Tetradron minimum (Tee), Scenedesmus obliquus (Sce), see Supplement 1 at www.int-res.com/articles/suppl/b017p197_supp.pdf for further information). Algae were grown using a modified WC medium (Guillard 1975). The concentrations of the most important nutrients for phytoplankton growth (50 μg P l−1, 1000 μg N l−1 and 1500 μg Si l−1) corresponded to a eutrophic lake with phosphorus limitation. The medium was buffered to maintain a pH of 7. Experimental communities were grown in semi-continuous culture using Erlenmeyer flasks (total culture volume [Vtot]; 152 ml). Every other day, 38 ml of Vtot was replaced with new medium (Vnew), resulting in a replacement rate (D) of 0.125 d−1 (D = Vnew/Vtot). Day:night cycle (16:8 h), light intensity (140 ± 5 μmol m−2 s−1 photosynthetic photon flux density) and experimental temperature (20 °C) were kept constant. A shaker-table (<100 rpm) kept the cultures in suspension. All culture work was done under sterile conditions.

The 2 stressors employed in our study were not meant to mimic naturally occurring disturbances but to impose 2 sources of mortality, potentially requiring different adaptations and thus allowing for trade-offs in species tolerances to either of the 2 stressors (see ‘Introduction’). pH reduction was a chemical stressor, whereas simulated grazing was a mechanical disturbance imposing strong size-selective mortality. For pH-reduction we used sterilised HCl and NaOH solutions (1 N). The pH was measured, reduced to pH 5, and on the following day it was set back to its original value. Since it was not possible to determine pH under sterile conditions, we prepared extra cultures for each experimental treatment that experienced pH reduction. These were cultured under the same conditions as the replicates of the treatments. To simulate grazing, we filtered 30% of the algal suspension through a 5 μm mesh-size membrane under sterile conditions. We chose this mesh size because it could be expected to affect small and single-celled strains to a lesser extent than larger and filamentous strains.

Experimental design and analyses

To test the effect of multiple stressors on phytoplankton communities of different diversity, we conducted 2 experiments. The monoculture experiment was designed to test how each single species in our species pool responded to the stressors pH reduction (pH) and grazing (gra). In the diversity experiment we exposed communities of different diversity and species composition to 2 stressors in different combinations (gra−pH, gra−pH, pH−gra, pH−pH).

In the monoculture experiment, each of the strains in our species pool was grown as single species culture. To gain an inoculum, the strains were grown as batch cultures under the experimental conditions described above for 5 d prior to the start of the experiment. For each strain, the same biovolume (1.82 × 106 μm3) was used to inoculate 3 replicate cultures per treatment (gra, pH, control; 11 × 3 × 3 = 99 experimental units). Duration of the monoculture experiment was 14 d. To account for an initial growth phase in the monoculture experiment, single stressors (either gra or pH) were applied on Day 7. Samples for microscopic analysis were taken before stressor application (Day 7), and on Days 8, 11 and 14. Water samples for phosphorus analysis were taken at the end of the experiment.

The diversity experiment consisted of multi-species cultures of different species richness in 2 scenarios. These were separated by the presence or absence of a ‘super-species’ with certain trait combinations for which we had clear expectations of its effect on the response to disturbance. Within the 2 scenarios, richness levels of the experimental communities were 2, 4 and 8 species. The lower richness levels were always nested subsets of the 8-species assemblage, as this was the only way to simulate the loss of richness while still keeping a certain best-performing species. It has been argued that nested subsetting restricts the test of species richness effects to certain species assemblages and thus does not represent a test of a general diversity–function relationship (Huston 1997). However, the alternative approach to random subsetting has also been questioned as it makes more speciose assemblages too similar in species composition compared to those which are less speciose, and this can strongly affect the variability of biomass production at different richness levels (Fukami 2001).
In Combination A, Sce, the species that (based on initial growth rate, competitive ability, biovolume losses and recovery rates) had been identified as the best-performing species in our monoculture experiment (see ‘Results: Monoculture experiment’ and Supplement 2 at www.int-res.com/articles/suppl/b017p197_supp.pdf), was present at each richness level, whereas the other species were randomly chosen from our species pool. The species in Combination B were a random selection from the pool after Sce had been excluded (2 species: Sce and Ooc [A] or Cry and Mel [B]; 4 species: Sce, Ooc, Tee and Chl [A] or Cry, Mel, Tes and Cyc [B]; 8 species: Sce, Ooc, Tee, Chl, Cry, Mel, Tes and Ped [A] or Cry, Cyc, Tee, Fra, Tes, Cry, Mel and Ped [B]). To obtain an inoculum, we grew each species in batch culture as described for the monoculture experiment. Multi-species cultures were inoculated with a biovolume of 3.41 × 10^8 μm^3, where the total inoculate biovolume consisted of equal fractions that corresponded to the number of species (2 species: 2 × [1.705 × 10^8 μm^3], 4 species: 4 × [0.853 × 10^8 μm^3], 8 species: 8 × [0.426 × 10^8 μm^3]; evenness = 1). Experimental communities were exposed to different stressor combinations (gra−gra, gra−pH, pH−gra, pH−pH) or remained undisturbed (control). Each treatment was replicated twice, which resulted in 60 (2 × 3 × 5 × 2) experimental units. The average duplication time of the species in our pool was 2 d during the initial growth phase (Days 0 to 7) of the monoculture experiment. To allow for 15 micro-algal generations, we chose 31 d for the duration of the experiment. This duration is shorter than the time span (35 to 50 d) in which the occurrence of competitive exclusion can be expected in phytoplankton communities cultured under semi-continuous conditions (Flöder & Burns 2005). It is likely, therefore, that the communities in our experiments were in a transitional stage (see Supplement 5 at www.int-res.com/articles/suppl/b017p197_supp.pdf for population dynamics). Experimental communities in the diversity experiment were exposed to the first disturbance on Day 7 and to the second disturbance on Day 13. To monitor the effect of the stressors on the communities, the experiment was sampled immediately before and after stressor application on Days 7, 8, 13 and 14. Further sampling was performed on Days 17, 27 and 31.

**Sample analyses**

Analysis of dissolved soluble reactive phosphate (SRP) followed the method by Strickland & Parsons (1978). In the monoculture experiment, the final concentration of SRP in control treatments was used as an indicator for competitive ability (comparable to Tilman’s [1982] limiting resource threshold R* in cultures at carrying capacity).

Algal abundance was analysed microscopically counting at least 400 cells. The dimensions of 20 individuals of each species were determined to calculate the average specific biovolume (Hillebrand et al. 1999). These data were used to calculate final total biovolume (B_tot), its temporal coefficient of variation (CVB) and the evenness index (E_B) at the end of the experiment:

\[
E_B = \frac{H_B'}{H_B'^\text{max}} \quad (1)
\]

\[
H_B' = -\sum p_i \ln(p_i) \quad (2)
\]

where \(H_B\) denotes the Shannon index, \(H_B'^\text{max}\) denotes the theoretical diversity maximum (= ln(species richness)) and \(p_i\) is the relative proportional contribution of species \(i\) to total biovolume (Washington 1984).

Growth rates were calculated based on biovolume according to:

\[
\mu = \frac{\ln(B_f) - \ln(B_i)}{t_2 - t_1} \quad (3)
\]

where \(\mu\) denotes the specific growth rate per day, \(t_1\) and \(t_2\) are 2 points in time during the course of the experiment and \(B_1\) and \(B_2\) denote the total community biovolume (diversity experiment) or the biovolume of 1 species (monoculture experiment) at \(t_1\) and \(t_2\), respectively.

To specifically analyse the resistance and resilience of the experimental communities, percent biovolume loss and recovery rate were calculated. In the monoculture experiment, biovolume in the disturbed treatments was expressed as a percentage of the average biovolume in the control for the respective strain. In the diversity experiment, biovolume in the disturbed treatments was expressed as a percentage of the biovolume in the control for the respective species composition, by which we randomly aligned the 2 replicates of treatment and control. Population and community losses due to disturbance were calculated for the day following the final disturbance (Day 14), and high resistance is reflected by low percentages of biovolume loss. The relative biovolume (i.e. in relation to the undisturbed control) was also employed to estimate recovery rates from specific growth rates after the final disturbance (Day 14). We calculated the rate of biovolume increase between Day 14 and the recovery date, which we defined as the day when the post-disturbance biovolume
reached the biovolume of undisturbed control cultures on Day 14. Exact recovery dates had to be calculated, since post-disturbance growth rates tended to change over time (see Supplement 3 at www.int-res.com/articles/suppl/b017p197_supp.pdf for details).

Statistical analyses

In the monoculture experiment, we tested the hypothesis that species differ in growth rate and percentage loss by a 1-factor ANOVA followed by a post hoc test (Tukey’s HSD). Species identity was the sole independent factor, though dependent variables consisted of growth rate and final remaining SRP, as well as the percentage of biovolume loss, the recovery rate and the recovery time for pH reduction and simulated grazing. Correlation analyses were used to test whether the recovery rates and the specific loss of biomass due to pH reduction and grazing were related.

In the diversity experiment, we performed separate factorial ANOVAs for Combinations A (Sce present) and B (Sce absent), as these combinations tested different hypotheses which would have been masked by a nested design (combination would have to be nested in richness) of a combined analysis. In Combination A, we tested whether the presence of Sce was sufficient to maintain resistance and resilience independent of the richness of the surrounding community. In Combination B, we tested whether more species were needed to maintain stability in the absence of a strongly performing species. The factorial ANOVA comprised richness and stressor regime as factors, and final biovolume, temporal biovolume variation, $E_B$, percent biovolume loss, recovery rate and co-tolerance index as response variables. The analysis of species’ co-tolerance to multiple stressors was based on the number of growing species recorded after the disturbances (see Supplements 4 & 5 at www.int-res.com/articles/suppl/b017p197_supp.pdf for details). For a diversity-independent measure, we subtracted the number of species that displayed positive growth rates after the application of Stressor 1 from the number of species that continued to grow after the application of Stressor 2. This difference was divided by the number of species recorded prior to the disturbances. Since the difference in the number of growing species can be either positive or negative, the resulting index of stressor-induced community co-tolerance (co-tolerance index) stretches from −1 to +1, where scores >0 indicate stressor-induced community tolerance. Scores between 0 and −0.5 indicate randomly distributed species co-tolerances, and scores below −0.5 point towards stressor-induced community sensitivity.

RESULTS

Monoculture experiment

In the monoculture experiment, we observed significant differences between species traits concerning competitive ability, as well as resistance and resilience in response to the stressors pH reduction and grazing. Initial growth rates ranged from 0.086 d$^{-1}$ (Tes) to 0.602 d$^{-1}$ (Sce). Sce, the fastest growing species, also displayed a high competitive ability by reducing the SRP concentration to a low level (3.3 μg l$^{-1}$). Biovolume losses due to pH reduction ranged from 1.52% (Ped) to 45.8% (Cry), and losses due to grazing ranged from 18.1% (Fra) to 49.5% (Tes). The species-specific losses in biovolume from pH reduction and grazing were uncorrelated ($r = 0.14$, $p = 0.68$, $N = 11$), whereas the recovery rates were positively correlated ($r = 0.83$, $p = 0.002$). Based on its high initial growth rate and competitive ability, the low biovolume loss due to pH reduction (17.0%) and grazing (23.5%) and the capability to recover within 5 d (recovery rate—pH: 0.038, gra: 0.053), we identified Sce as the best-performing species in our species pool (see Supplement 2 for detailed results).

Community biovolume and evenness in the diversity experiment

In the diversity experiment, the factors species richness and stressor regime affected $B_{tot}$ in both species combinations, but at least partly with opposing signs. In Combination A, final $B_{tot}$ decreased with species richness (Table 1, Fig. 1), whereas the stressor regime had no significant effect on $B_{tot}$. Thus, the presence of Sce resulted in non-significant stressor effects, but this effect of a high-performing species was diluted at higher species richness. Correspondingly, we found significant stressor × richness interaction, because the negative effect of richness on $B_{tot}$ was found in disturbed treatments only. In Combination B, increasing species richness increased $B_{tot}$ in the control. $B_{tot}$ was significantly reduced in comparison to the control when grazing (gra−gra and pH−gra) was the second stressor (Tukey’s HSD). When pH reduction was applied as
the second stressor (pH–pH and gra–pH) treatment, \( B_{\text{tot}} \) did not differ from the control. Significant stressor × richness interaction showed that the increase in \( B_{\text{tot}} \) was restricted to the control and those stressor regimes reducing \( B_{\text{tot}} \) (gra–gra and pH–gra), whereas richness reduced \( B_{\text{tot}} \) in the stressor combination pH–pH.

Independent of the multiple stressor scenarios applied, disturbance always significantly increased the temporal variability of biovolume in both species combinations (Fig. 1, Table 1). Control treatments showed the lowest \( \text{CV}_B \) throughout. In Combination A, \( \text{CV}_B \) increased with species richness (significant main effect; Table 1); thus, more species-rich assemblages showed higher variation of biovolume through time. In Combination B, we found a significant interaction between richness and stressor regime which reflected a strong positive richness effect on \( \text{CV}_B \) only for the gra–pH combination. For none of the stressor regimes did we observe a reduction in variability with richness.

### Table 1. Factorial ANOVA results for species combinations A and B: effects of stressor regime (stress) and species richness on biovolume (\( B_{\text{tot}} \)), coefficient of variation (\( \text{CV}_B \)) and biovolume evenness (\( E_B \)) at the end of the experiment. Degrees of freedom in the error term were 15. \( B_{\text{tot}} \) data of Combination B failed to become homoscedastic. Since data transformation increased the level of heteroscedasticity, we used untransformed data for the analysis.

<table>
<thead>
<tr>
<th>Effect</th>
<th>df</th>
<th>( B_{\text{tot}} ) F</th>
<th>( B_{\text{tot}} ) p</th>
<th>( \text{CV}_B ) F</th>
<th>( \text{CV}_B ) p</th>
<th>( E_B ) F</th>
<th>( E_B ) p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Combination A</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Richness</td>
<td>2</td>
<td>15.16 0.0003</td>
<td>8.688 0.0031</td>
<td>109.9 &lt;0.0001</td>
<td></td>
<td></td>
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<tr>
<td>Stress</td>
<td>4</td>
<td>0.510 0.7293</td>
<td>15.50 &lt;0.0001</td>
<td>8.133 0.0011</td>
<td></td>
<td></td>
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<tr>
<td>Stress × Richness</td>
<td>8</td>
<td>3.122 0.0274</td>
<td>2.387 0.0697</td>
<td>4.038 0.0097</td>
<td></td>
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<tr>
<td><strong>Combination B</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Richness</td>
<td>2</td>
<td>5.525 0.0159</td>
<td>1.179 0.3347</td>
<td>58.20 &lt;0.0001</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stress</td>
<td>4</td>
<td>32.28 &lt;0.0001</td>
<td>11.82 0.0002</td>
<td>51.26 &lt;0.0001</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stress × Richness</td>
<td>8</td>
<td>3.556 0.0165</td>
<td>6.181 0.0013</td>
<td>34.59 &lt;0.0001</td>
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</tbody>
</table>

Fig. 1. Total biovolume (\( B_{\text{tot}} \)) at the end of the diversity experiment, temporal coefficient of biovolume variation (\( \text{CV}_B \)) and final biovolume-based evenness (\( E_B \)) in species combinations A (●) and B (○) in different combinations of 2 stressors (grazing [gra] and pH); 2, 4, and 8 species (Sp) denote levels of species richness being tested. 
Species richness explained 90% of the variance in $E_B$ in Combination A (Fig. 1, Table 1). $E_B$ at the end of the experiment was lowest for 2 species, highest for 4 species and intermediate in treatments containing 8 species. The stressor regime affected $E_B$ levels, but not its unimodal response to richness. In Combination B, $E_B$ decreased with increasing richness (significant main effect; Table 1), but with strong interaction with the stressor regime. $E_B$ increased with richness under the stressor regimes pH–pH and gra–pH, but decreased in the control and under gra–gra and pH–gra disturbance regimes (significant 2-way interaction; Fig. 1, Table 1).

**Community resistance, resilience and co-tolerance in the diversity experiment**

Species richness and stressor regime affected percent biovolume loss in both species combinations (Fig. 2). In both A and B, high biovolume loss was recorded for the stressor regime gra–pH, whereas the lowest percentage of biovolume was lost under the opposite regime, pH–gra. This result indicates that not only the type of disturbance but also its sequence is highly important for its effect. In Combination A, the percentage of biovolume loss decreased with richness (significant main effect; Table 2), with 4 and 8 species levels losing less biovolume due to disturbance than 2 species levels. The opposite result was found for Combination B, where 4 and 8 species levels lost more biovolume than 2 species levels (significant main effect; Table 2). Diverging from the overall trend, percent biovolume loss in Combination B did not increase with richness under the stressor regime gra–gra (significant 2-way interaction; Table 2).

<table>
<thead>
<tr>
<th>Effect</th>
<th>df</th>
<th>Biovolume loss (%)</th>
<th>Recovery rate</th>
<th>Co-tolerance</th>
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<tr>
<td></td>
<td></td>
<td>$F$</td>
<td>$p$</td>
<td>$F$</td>
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<tr>
<td><strong>Combination A</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Richness</td>
<td>2</td>
<td>24.46</td>
<td>0.0001</td>
<td>0.573</td>
</tr>
<tr>
<td>Stress</td>
<td>3</td>
<td>22.39</td>
<td>&lt;0.0001</td>
<td>44.14</td>
</tr>
<tr>
<td>Stress $\times$ Richness</td>
<td>6</td>
<td>1.918</td>
<td>0.1587</td>
<td>4.004</td>
</tr>
<tr>
<td><strong>Combination B</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Richness</td>
<td>2</td>
<td>9.396</td>
<td>0.0035</td>
<td>1.211</td>
</tr>
<tr>
<td>Stress</td>
<td>3</td>
<td>26.19</td>
<td>&lt;0.0001</td>
<td>22.60</td>
</tr>
<tr>
<td>Stress $\times$ Richness</td>
<td>6</td>
<td>5.170</td>
<td>0.0077</td>
<td>11.13</td>
</tr>
</tbody>
</table>
The stressor regime affected recovery rates in both species combinations (Fig. 2, Table 2). Stressor regimes leading to strong biovolume reduction tended to show the highest recovery rates (gra−pH in A and B; pH−pH in A; Fig. 2). Richness had no significant main effect, but we found significant richness × stressor interactions in both combinations (Table 2). In Combination A, the recovery rate decreased with richness in the gra−pH regime, whereas, under the other disturbance regimes, it tended to increase with richness or remained unchanged (Fig. 2). Recovery rates in Combination B decreased with richness under stressor regimes gra−pH and pH−pH, but increased with richness under the pH−gra regime.

The co-tolerance index indicated stressor-induced community tolerance in all treatments of Combination A (2 and 8 species levels: score > 0; 4 species level: score ~ 0) (Fig. 3). The index was lowest at intermediate richness (significant main effect; Table 2) and strongly decreased after the second disturbance in the 2 species level of the pH−pH regime (significant richness × stressor interaction; Table 2). In Combination B, the co-tolerance index increased with increasing species richness (significant main effect; Table 2) because the average score at the 2 species level (less than −0.5) pointed towards stressor-induced community sensitivity, whereas average scores at the 4 and 8 species levels were just below zero, indicating randomly distributed co-tolerances.

**Species-specific responses in the diversity experiment**

In Combination A, Sce dominated B_{tot} of all experimental communities (2, 4, and 8 species levels) throughout the entire experiment (see Supplements 6 & 7 at www.int-res.com/articles/suppl/b017p197_supp.pdf for details). Higher dominance of Sce was correlated to higher B_{tot}, i.e. higher evenness in the presence of a high-performing species resulted in lower biomass production (Fig. 4A).

The dominance structure was much more variable in Combination B, differing among species levels and with stressor regime (see Supplements 6 & 7 for details).
details). At the end of the experiment, *Mel* and *Cry* co-dominated the 2 species level of the controls, *Mel* dominated the 4 species level with *Cry* being abundant and *Cyl* was highly dominant at the 8 species level. Prior to stressor application, *Mel* dominated at the 2 and 4 species levels, and *Fra* contributed most of *Btot* at the 8 species level. After the application of the stressor regimes, a complex pattern between richness and the stressor regimes appeared. In the 2 and 4 species levels, *Mel* remained dominant after pH−pH and gra−pH disturbance sequences. Under pH−gra and gra−gra regimes, *Cry* became dominant in the 2 species level, and *Cyc* became dominant under the gra−gra regime in the 4 species level. The 8 species level shifted from *Fra* dominance to *Cyl* under the gra−gra and pH−gra regimes, whereas *Fra* retained dominance under the pH−pH and gra−pH regimes. In Combination B, evenness was also negatively correlated to *Btot*, but, in the absence of a high-performing species, biomass production was low (Fig. 4B).

**DISCUSSION**

**Species performance in single and mixed cultures**

The monoculture experiment revealed species-specific differences concerning the traits initial growth rate, competitive ability, as well as biovolume loss and recovery rate in response to the stressors pH reduction and grazing. Although some species were functionally similar with regard to biovolume production (indicating redundancy sensu Walker 1992), these species differed in resource use, biovolume loss and recovery rate. Thus, trait variance in the diversity experiment was certain to increase with increasing species richness.

*Sce* was identified as the best-performing species. It had the highest initial growth rate, a high competitive ability for limiting nutrient phosphorus, low biovolume losses in response to the stressors pH reduction and grazing, and it recovered quickly from the disturbance caused by those stressors. This species was included in the inoculum of each richness level of Combination A and was absent in Combination B. In the diversity experiment, *Sce* became dominant regardless of the richness level of the experimental communities. Consequently, *Eg* in Combination A was generally low, but it increased with increasing species richness. This was mainly due to *Chl*, another productive species that became subdominant at higher richness levels in Combination A.

While *Sce* performed as expected in mixed culture, the performance of other species in mixed culture differed from their performance in the monoculture experiments. *Chl* for example became subdominant at the 2 and 4 species levels, although *Ped* had displayed lower losses and higher recovery rates in response to single stressors in the monoculture experiment. Such discrepancies in species responses to single and multiple stressors are likely to be due to interactions that cause non-additive effects (Breitburg et al. 1998, Vinebrooke et al. 2004). In the communities of Combination B, *Mel* performed better at the 2 species level than at the 4 and 8 species levels, indicating that a species’ ability to tolerate multiple stressors can depend on the competitive surrounding (Vinebrooke et al. 2004).

Moreover, not only the type of stressor but also the temporal sequences affected the response, as mixtures in both species combinations responded differently to pH−gra compared to gra−pH sequences. Previously, Fukami (2001) found that community composition diverges with different sequences of the same type and intensity of disturbance. Obviously, the ability to cope with a certain stressor depends strongly on the disturbance history, which lends support to the idea of community co-tolerance or sensitivity, i.e. the change in a community’s susceptibility to disturbance by shifts in community composition (Vinebrooke et al. 2004).

**Final biovolume and temporal variation in biovolume**

Both artificial disturbances were highly effective, as we observed up to 50% loss of biovolume in the monocultures (Supplement 2) and an average of 35% loss of biovolume after pH reduction and 37% loss after grazing applications in the diversity experiment. Indeed, both stressors increased the temporal variation of total biovolume compared to the control. The monoculture experiment showed that species’ responses to the different disturbances was uncorrelated in our species pool.

In Combination A, community performance depended strongly on *Sce* alone, and any decrease in the relative proportion of this species reduced biomass production and stability. Thus, adding more species reduced stability, and higher evenness was related to lower *Btot*. In fact, the presence of *Sce* inhibited the presence of long-lasting stressor effects on *Btot*. Thus, Combination A corresponds well to the model settings of Norberg et al.’s (2001) study of the relationship between trait variance and productivity.
in stable and variable environments. Contradicting prevalent ecosystem theory, according to which niche differentiation increases production in stable environments (Tilman et al. 1997, Hector et al. 1999), Nörberg et al. (2001) argued that a single species with certain optimal traits allows the highest productivity under stable environmental conditions. An increase in trait variance reduces productivity, because species are present that perform suboptimally under the given environmental conditions. The model predicts that long-term productivity in variable environments increases with trait variance, because species are present that are able to respond to environmental changes. However, in our experiment, the most productive species was also the one species promoting stability, thereby preventing that an increase in trait variance with higher species richness increased biovolume production in the disturbed treatments.

Communities without Sce produced less biomass, and higher $B_{tot}$ was observed in Combination A than in B across all richness levels. Given our experimental design, we cannot generalise beyond the compositions tested here. However, the importance of species identity effects in biodiversity–production relationships is frequently found and often related to trait combinations (for a discussion see Hillebrand & Matthiessen 2009). The presence of such ‘super-species’ that show high performance for multiple situations is a frequent phenomenon in experiments with a small species pool and low complexity of physical or biological structure (Weis et al. 2008, Ptacnik et al. 2010).

When communities without this high-performing species (Combination B) were exposed to multiple stressors, there was no consistent relationship of $B_{tot}$ and $E_{B}$ to the different levels of species richness. $B_{tot}$ increased and $E_{B}$ decreased with species richness under those stressor regimes that were connected with an overall low final biovolume and high biovolume variability (gra–gra and pH–gra). Cyl dominated at the highest level of species richness in these treatments. When multiple stressors decreased $B_{tot}$ only marginally compared to the control, $B_{tot}$ decreased (pH–pH) and $E_{B}$ increased (pH–pH, gra–pH) with species richness. Here Fra dominated at the highest richness level. Increasing richness (i.e. trait variance) increased the biovolume production of communities only in certain stressor combinations, a situation which was caused by local dominance rather than by niche separation or facilitation, since high levels of $B_{tot}$ in species-rich treatments were connected with low levels of $E_{B}$ and vice versa (Doak et al. 1998, Cottingham et al. 2001). This emphasises the importance of addressing not only richness, but also species identity and dominance when investigating biodiversity effects in general (Wilsey et al. 2005, Hillebrand et al. 2008), and DSH especially. Moreover, our results indicate that the effect of response diversity is context dependent, as the positive effect of richness on $B_{tot}$ depends on the stressor regime.

Richness did not reduce $CV_B$ in either Combination A or B. By contrast, $CV_B$ increased with richness in A and in some stressor sequences of B. The effect of richness on stability was thus not expressed as a buffering of the temporal variability but rather by providing fast recovery (see below). However, the presence of Sce reduced $CV_B$ (cf. Combinations A and B in Fig. 1); thus, the presence of a high-performing species maintained low variability. Similarly, Steiner et al. (2005) found the dominance of fast-growing green algae (Ankistrodesmus sp. and Chlorella sp.) to be negatively related to community biovolume variability.

### Resistance and resilience to multiple stressors

In Combination B biovolume loss tended to be higher and the recovery rate lower than in A. Resistance increased with increasing species richness when a high-performing species was present at all richness levels (Combination A), whereas the opposite was true in the absence of a high-performing species (Combination B). When Sce was present, it remained dominant under the impact of disturbance and carried the post-disturbance production. At higher richness levels, sub-dominant and abundant species also continued to grow, explaining the positive richness effect in Combination A. Similar to natural systems (O’Neill 1999), resistance to multiple stressors in our experiment was a function of immediate losses to both stressors, and, since the stressors were applied with a 7 d time lag, recovery from the stressor that was applied first. Highest losses were connected to communities where dominant, fast-growing species after the first stress event either stopped growing or continued to grow at only a very low rate. Compensatory growth could be observed in species that had performed poorly before the first stressor was applied, and thus had started out from a comparatively low specific biovolume.

Regardless of the differences in dominance structure development, we found evidence for stressor-induced community tolerance in both species combinations. The co-tolerance index was high in the presence of Sce, whereas, though low, it increased with species richness in the absence of Sce. Thus, species
traits determining the tolerance to one stressor can influence how communities respond to other stressors (Vinebrooke et al. 2004). Stressor-induced species tolerances result in antagonistic community responses to multiple stressors and can explain the tendency towards the antagonistic multiple stressor effects often found in autotrophic communities (Vinebrooke et al. 2004, Crain et al. 2008).

Resilience was more strongly affected by the stressor regime than by species richness, which showed no significant main effect. The recovery rates tended to be higher in treatments that had experienced high losses in both species combinations. A potential explanation for this pattern is a ‘self-regulation’ of the steady state, which is one of the characteristics of the semi-continuous culture technique that we used in our experiments (Monod 1950). In the absence of disturbance, loss and growth rates equal zero at steady state. If disturbance reduces total biovolume, the resulting reduction in resource use causes resource concentrations to increase, which initiates the recovery of the community. The more biovolume is removed, the more resources become available and the higher the recovery rate (Flöder & Sommer 2006).

The effect of richness on resilience was embedded in significant richness × stressor interactions in both species combinations. With Sce, recovery rates increased with species richness under those stressor regimes that led to small or moderate losses, whereas they decreased with richness when biovolume loss was high. The latter was presumably connected to higher resource availability, allowing Sce to respond with high growth rates leading to faster recovery—an effect dilated by more species being present. When biomass losses were low, recovery was driven by more species showing compensatory growth, leading to increased resilience with increased richness. This interaction corresponds to the results of a study on diversity effects on stability of multitrophic communities (Steiner et al. 2005, 2006), where resilience was positively related to diversity at low productivity, but not at high productivity. Increasing resilience was related to the increasing dominance of fast-growing species of small green algae. In randomly assembled communities, resilience was related to the functional trait of the species present. When the same species that had dominated before the stressors were applied dominated at the end of the experiment, recovery rate decreased with richness. When the dominance structure changed continuously and the dominant species did not recover from the multiple stressors, recovery rates were either low at all richness levels or increased with species richness.

Conclusions

Both stressors, pH reduction and grazing, play important roles in natural systems. Episodic acid stress threatens freshwater ecosystems in acidified regions of Europe and northern America, because phases of droughts and floods are expected to intensify due to climatic change. Drawdown of the water table in water-logged organic soils cause previously reduced compounds to become re-oxidised, which results in pulses of acidic, sulphate-rich water (Devito et al. 1999, Laudon et al. 2004). The artificial grazing that we employed in our experiments consisted of a rather unspecific removal of microalgae. In natural systems grazing can affect phytoplankton communities in different ways. Grazers can be general in their feeding behaviour, or they can be specialised feeders (Hillebrand & Shurin 2005). The rapidly increasing grazing pressure of a growing Daphnia spp. population on small-celled phytoplankton species in spring, resulting in the clear water stage of northern temperate lakes, is a well researched phenomenon (e.g. Sommer et al. 1986). Since Sce is prone to grazing by Daphnia spp., this type of grazing would have affected the ranking of species performance and community development in our study. This fact emphasises that the concept of super-species depends on species traits and stressor identity. ‘Super-species’ occur in several phytoplankton biodiversity-ecosystem functioning experiments, especially under highly controlled culture conditions (Placnik et al. 2010). Natural fluctuations in real ecosystems will potentially reduce the role of such species, as the increased potential niche space allows for a variety of successful trait combinations rather than only one. Isbell et al. (2011) showed that in fact more species contribute to the ecosystem functioning of environments that change in space or time than those in constant environments.

The interdependence of stressor regimes, species traits and species richness determines which mechanisms stabilise natural communities. If dominant species remain the best performers regardless of disturbance, stability will depend on the population dynamics of species with these successful functional traits. If, on the other hand, disturbance or environmental change reverses the hierarchy of successful functional traits and dominant species become rare or lost (Jablonski 1994, Grime 1998), compensatory growth of rare species occurs. In natural systems, such an effect could insure communities against complete failure.
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LITERATURE CITED


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