# 1 Selection in response to community diversity alters plant performance and

# 2 functional traits

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#### 24 ABSTRACT

25 In grassland biodiversity experiments the positive biodiversity-ecosystem functioning 26 relationship generally increases over time. However, we know little about the underlying 27 short-term evolutionary processes. Using five plant species selected for twelve years in a 28 biodiversity experiment in mixture or monoculture and plants without such a selection 29 history, we assessed whether differential selection altered productivity, biodiversity effects, 30 and functional trait differences within newly assembled monocultures and 2-species 31 mixtures. Plants without past community selection history produced the lowest assemblage 32 biomass and showed the weakest biodiversity effects. In newly assembled mixtures, plants 33 with a selection history in mixtures produced more biomass than plants with a monoculture 34 selection history. Biodiversity effects were generally positive and differed significantly 35 between selection histories. However, contrary to our expectations, biodiversity effects were 36 not stronger for mixture-type plants. Biodiversity effects were influenced by both trait 37 differences between plants and community-weighted means, but these relationships were 38 mostly independent of selection history. Our findings suggest that twelve years of selection 39 history in monocultures or species mixtures differentiated plants of each species into 40 monoculture- and mixture-types. Such rapid evolution of different community-types within 41 grassland species and its effect on ecosystem services and functioning are likely to be 42 important for species conservation practice.

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#### 44 **1. Introduction**

The importance of biodiversity for maintaining ecosystem functions such as productivity or nutrient cycling is now well established (Cardinale et al., 2012). The biodiversity–productivity relationship is positive in grassland ecosystems (e.g. Isbell et al., 2011; Tilman et al., 2001), with biodiversity increasing multiple ecosystem functions (Soliveres et al., 2016). The positive effect of biodiversity strengthens with time (Cardinale et al., 2007; Reich et al., 2012), suggesting that complementarity between the co-occurring species can increase over time (Fargione et al., 2007).

52 Despite more than a decade of research on the biodiversity-productivity relationship 53 (e.g. Reich et al., 2012), we know little about the evolutionary mechanisms that potentially 54 affect species interactions (Thorpe et al., 2011). Selection acting on traits may increase 55 ecological combining ability (Aarssen, 1983; Harper, 1977) via niche differentiation in plant 56 mixtures (Zuppinger-Dingley et al., 2014). Such adaptation occurs when there is either 57 sufficient standing genetic variation in a population and a sorting out of the most suitable 58 genotypes (Fakheran et al., 2010) or by recombination and novel mutations (Anderson et al., 59 2011). Furthermore, plants may adapt to a novel environment by phenotypic plasticity (Price 60 et al., 2003; Turcotte and Levine, 2016), thus changing their morphology without genotypic 61 changes. Epigenetic variation may contribute to phenotypic variation and thus provides a 62 further source for selection and adaptation (Bossdorf et al., 2008).

Whereas the influence of environmental factors on adaptive responses of plant populations is well studied (e.g., Joshi et al., 2001; Schmid, 1985), much less effort has been devoted to studying the influence of community diversity on a species' performance (but see Kleynhans et al., 2016; Lipowsky et al., 2011). Based on previous observations in experimental ecosystems suggesting a "division of labor" among species in plant mixtures, it is likely that community diversity plays a role in the evolution of plant functional trait

69 variation. For example, in forests more diverse tree communities express greater crown 70 complementarity (Niklaus et al., 2017; Williams et al., 2017). In diverse grassland 71 communities increased complementarity effects, as estimated by the additive partitioning 72 method of Loreau and Hector (2001), promote community productivity via diversification of 73 the canopy structure and hence light and space use (Allan et al., 2011; Spehn et al., 2000), 74 soil resource partitioning (Fornara and Tilman, 2008; Roscher et al., 2008; von Felten et al., 75 2009), root depth distribution (Mueller et al., 2013) and distribution of leaf mass (Wacker et 76 al., 2009). However, these *in situ* observations do not allow separating the contribution of 77 phenotypic plasticity from potential underlying shifts in the population structure due to 78 selection of different community-types. To understand to which extent evolutionary 79 processes drive these differences, it is thus important to assess biodiversity effects in a 80 common environment.

81 Using the additive partitioning method (Loreau and Hector, 2001), the net 82 biodiversity effect (NE) can be partitioned into complementarity (CE) and sampling effects 83 (SE). When the CE drives over-yielding, most species are expected to contribute to higher 84 productivity in more diverse communities. In contrast, when the SE drives over-yielding, a 85 few dominant species increase community productivity in species mixtures. The CE is 86 therefore related to coexistence and trait variation between species as it inherently suggests a 87 differentiation in functional traits (Cadotte et al., 2009; Flynn et al., 2011). Conversely, the 88 SE should be driven by traits of the dominant species and thus by community-weighted trait 89 means (CWMs); an increase in CWMs (e.g., taller plants) should increase biodiversity 90 effects (Cadotte, 2017; Roscher et al., 2012).

91 The use of functional traits to define species' niches has a long history in
92 evolutionary ecology (Roughgarden, 1974; Schoener and Gorman, 1968; van Valen, 1965).
93 Recently it has become a popular approach in functional ecology (Violle et al., 2007) to

explain mechanisms of species coexistence and ecosystem functioning (Hart et al., 2016;
Kraft et al., 2015). However, we do not know how such trait-based niches and the associated
functional traits may evolve (Roscher et al., 2015; Sterck et al., 2011).

97 We tested whether community diversity can act as a selective environment shaping 98 biodiversity effects and functional traits. To test for the heritability of these effects and traits, 99 we grew offspring of plants grown for twelve years in a biodiversity experiment in 100 monocultures and two-species mixtures in a common environment. We measured biomass 101 production and traits of individual plants in monocultures and mixtures established with 102 seedlings from either a selection history of experimental monoculture or mixture 103 communities in a biodiversity field experiment (Jena Experiment, see Roscher et al., (2004) 104 for details) or in monoculture fields from the commercial seed supplier which provided the 105 original seeds for the Jena Experiment in 2002. We refer to the plants growing in Jena since 106 2002 in mixture or monoculture as mixture- and monoculture-type plants, respectively. The 107 plants derived from seeds obtained from the commercial supplier in 2014 are referred to as 108 naïve plants.

109 We previously assessed selection outcomes in the Jena Experiment after eight years 110 and one controlled sexual reproduction cycle (Zuppinger-Dingley et al., 2014). Here, we 111 prolonged the selection treatment by four more years, added a second controlled sexual 112 reproduction cycle, and refined our insights by measuring all individuals in test communities, 113 thus also allowing us to assess intra-specific variation within communities. We included 114 naïve plants as a control treatment without selection. We hypothesized that during the twelve 115 years of selection in the experimental field, mixture-type plants may have evolved increased 116 mixture performance. In turn, this may be associated with a larger NE, in particular CE, and 117 larger between-species trait variations. Conversely, we expected monoculture-type plants to 118 have evolved increased monoculture performance, which should be related to larger within119 species trait variation.

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- 121 **2. Materials and methods**
- 122 2.1 Plant selection histories

123 To test whether plant types selected over twelve years in mixtures outperform those 124 types selected in monocultures when assembled in mixture test communities, we chose five 125 species grown in monoculture and mixture plots in the Jena Experiment (Jena, Thuringia, 126 Germany, 51°N, 11°E, 135 m a.s.l., see Roscher et al., (2004) for experimental details): 127 Plantago lanceolata L., Prunella vulgaris L., Veronica chamaedrys L., Galium mollugo L. 128 and Lathyrus pratensis L. The study species had previously been classified into the following 129 functional groups (Roscher et al. 2004): small herbs (V. chamaedrys, P. vulgaris, and P. 130 *lanceolate*), tall herb (G. mollugo) and legumes (L. pratensis). Plants with Jena Experiment 131 selection history were sown in either mixture or monoculture in 2002 (selection history 132 "mixture" and "monoculture", respectively) and had undergone twelve years of selection 133 from 2002 until 2014 in either plant monocultures or species mixtures. The species 134 compositions in the experimental plots in Jena were maintained by weeding three times per 135 year in spring, summer and autumn and by mowing twice per year at peak biomass in spring 136 and summer.

We used plant progeny from three different selection histories for the experiment.
Plants without selection history in the Jena Experiment (selection history "naïve") were
obtained from a commercial seed supplier (Rieger Hoffmann GmbH, BlaufeldenRaboldshausen, Germany).

### 142 2.2 First controlled seed production

143 In spring 2010, plant communities of 48 plots (12 monocultures, 12 two-species 144 mixtures, 12 four-species mixtures and 12 eight-species mixtures) of the Jena Experiment 145 (Roscher et al., 2004) were collected as cuttings and transplanted into an experimental 146 garden in slug-exclosure compartments at the University of Zurich, Switzerland Switzerland 147 (47°33'N, °37'E, 534 m a.s.l.), in the identical plant composition as the original experimental 148 plots for the first controlled sexual reproduction among co-selected plants (Zuppinger-149 Dingley et al., 2014). In spring 2011, the seedlings produced from the seeds of the first 150 controlled sexual reproduction in Zurich were transplanted back into those plots of the Jena 151 Experiment from where the parents had originally been collected. In these newly established 152 plots, plant communities with an identical composition to the original communities were 153 maintained for three years until 2014.

154

#### 155 2.3 Second controlled seed production

156 To ensure a second sexual reproductive event for the collection of seed material, entire 157 plant communities from the experimental plots replanted in Jena in 2011 were excavated in 158 March 2014. As for the first controlled seed production, the plants from Jena were used to 159 establish plots with an identical plant composition in the experimental garden at the 160 University of Zurich,. We added a 30 cm layer of soil (1:1 mixture of garden compost and 161 field soil, pH 7.4, commercial name Gartenhumus, RICOTER Erdaufbereitung AG, Aarberg, 162 Switzerland), to each plot to ensure the plants established well. Mesh fabric netting around 163 each plot minimized the possibility of cross-pollination between the same species from 164 different experimental plots. We collected seeds from seven monoculture plots, one 4-165 species mixture plot and six 8-species mixture plots in the experimental garden. We did not 166 include seeds from 2-species mixtures as we expected that the community diversity selection

pressure may not be different enough from monocultures. Seeds from different mother plants were pooled together and cleaned manually for three species and mechanically for two species (*P. lanceolata* and *P. vulgaris*). The dry seeds were stored for 5 months at 5° C for cold stratification.

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172 2.4 Experimental set up

173 All seeds were germinated in germination soil ("Anzuchterde", Ökohum, 174 Herbertingen, Germany) under constant conditions in the glasshouse without additional light 175 in December 2014. From 25 February to 13 March 2015, seedlings were planted in 176 monocultures of four individuals or 2-species mixtures of two plus two individuals into pots 177 (two liters) filled with neutral agricultural soil (50% agricultural sugar beet soil, 25% perlite, 178 25% sand; Ricoter AG, Aarberg, Switzerland). At the beginning of the experiment the 179 studied plants were infested by fungus gnats (Bradysia spp.), which caused seedling 180 mortality during the experiment. Seedlings that died in the first two weeks were replaced 181 with seedlings of the same species and age.

182 We planted species assemblages in six blocks (replicates); each block included the full 183 experimental design. Species pairs were chosen according to seedling availability. The full 184 design (every possible species combination) was intended but could not be realized due to 185 seedling mortality and low germination rates for some species (e.g. G. mollugo). Within each 186 block, pots were placed on three different tables in the glasshouse in a randomized fashion 187 without reference to selection history or assembly treatment. During the timeframe of the 188 experiment we did not move pots but noted their position in the glasshouse. Single pots 189 always contained four plants of the same selection history. Each selection history  $\times$  species 190 assembly combination was replicated five to six times depending on plant availability. We 191 planted 30 monoculture and 42 mixture assemblages with mixture selection history, 30 monoculture and 60 mixture assemblages with monoculture selection history and 24
monoculture and 35 mixture assemblages with naïve selection history, a total of 221 pots and
884 plants (Appendix S2 for monoculture identities and species combinations).

195 During the experiment, the plants were initially kept at day temperatures of  $17-20^{\circ}C$ 196 and night temperatures of 13-17°C without supplemental light. To compensate for 197 overheating in summer, an adiabatic cooling system (Airwatech; Bern, Switzerland) was 198 used to keep inside temperatures constant with outside air temperatures. The plants were not 199 fertilized. Due to an infestation of white flies (*Trialeurodes vaporariorum*, Westwood 1856) 200 and spider mites (Tetranychidae spp., Donnadieu 1875), we applied the insecticide 201 SanoPlant Neem (1% Azadirachtin A (10 g/l); Maag AG) three times. The fungicide Fenicur 202 (Oleum foeniculi, Andermatt Biocontrol) against powdery mildew (Podosphaera spp.) was 203 applied twice. Plant height, leaf thickness, specific leaf area (SLA) and individual 204 above ground biomass were measured after twelve weeks of the experiment from 18 May to 4 205 June 2015. Leaf thickness was measured for three representative leaves using a thickness 206 gauge. Specific leaf area (SLA) of up to 20 representative leaves (depending on the leaf size 207 of the species) of each species in a pot was measured by scanning fresh leaves with a Li-208 3100 Area Meter (Li-cor Inc., Lincoln, Nebraska, USA) immediately after harvest and 209 determining the mass of the same leaves after drying. Plant height and individual 210 aboveground biomass were measured a second time after 24 weeks from 18–25 August 2015 211 at the end of the experiment. All four individuals in a pot were sampled. Research assistants, 212 who were not informed of the specific experimental treatments, assisted in the regular 213 measurements and harvesting of plants at the end of the experiment.

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#### 217 2.5 Data analysis

218 We calculated pot-wise aboveground community biomass (plant community 219 production) as the sum of the biomass of the four individual plants. Because the first measure 220 assessed growth and the second regrowth, the harvests were analyzed separately. Relative 221 between-species differences (RDs, absolute difference between two species divided by the 222 mean of the two) in plant height (first and second harvest), leaf thickness (first harvest) and 223 SLA (first harvest) were calculated for mixture assemblages. Relative differences within 224 species were calculated for both mixture and monoculture assemblages taking the relative 225 difference between two individuals of the same species per pot. SLA outliers (> 99% 226 percentile) were replaced with a maximum value (the 99% percentile, n = 6). Furthermore, 227 we calculated community-weighted means (CWMs) and pot standard deviation (SDs) for the 228 same traits (R package "FD", Laliberté and Legendre 2010; Laliberté et al., 2014). Pots with 229 dead plant individuals were excluded from the calculation of community-weighted means, 230 but were included for the other data analyses. Net biodiversity effects (NE) were calculated 231 by comparing the 2-species mixtures with the average monoculture and partitioned according 232 to Loreau and Hector (2001) into complementary (CE) and sampling (selection) effects (SE). 233 This partitioning approach allows assessing how CE and SE contribute to the NE (Loreau 234 and Hector 2001). To avoid confusion with the term selection used for the selection-history 235 treatment, we here use the term "sampling effect" for the SE (see Zuppinger-Dingley et al., 236 2014). Additive partitioning calculations were based on the difference between the observed 237 yield of each species in the mixture and the monoculture yield for that species and selection 238 history averaged across blocks. Absolute values of CE and SE were square root-transformed 239 and the original signs put back on the transformed values for analysis (Loreau and Hector 240 2001). Differences in these measures between mixtures assembled from plants with 241 monoculture selection history and mixtures assembled from plants with mixture selection history would suggest differential evolution of trait-based niches between species as apotential mechanism underlying biodiversity effects.

244 All statistical analyses were done in R (Version 3.2.3, R Core team 2016). Mixed-245 model analysis was done using the R-package asreml (VSNI international, 2016) and results 246 were assembled in ANOVA tables. Selection-history treatment (naïve, monoculture, 247 mixture), assembly treatment (monoculture vs. 2-species mixture assemblages), species 248 identity of monoculture assemblages and of mixture assemblages (in short "species 249 assembly") and interactions of these were fixed-effects terms in the model. Table (including 250 blocks) was a random-effects term in the model. CWMs, RDs, within-species differences 251 and SDs of plant height, SLA and leaf thickness were added as covariates to determine the 252 influence of these covariates on community biomass and biodiversity effects.

253

## 254 **3. Results**

# 255 *3.1 Plant selection history and community productivity*

Assemblages consisting of plants with naïve selection history produced the lowest community biomass at both harvests (Fig. 1; Table 1). At the second harvest, such higher productivity of selected plants was stronger in 2-species mixtures than in monocultures (Fig. 1; Table 1). At the second harvest, mixture-type plant assemblages outperformed monoculture-type plant assemblages and this effect was marginally more pronounced in the 2-species mixtures (Fig. 1; Table 1).

Species identity in monoculture or mixture assemblages strongly influenced community productivity and, especially at the first harvest, the interaction terms with selection history were significant (Table 1). For example, at the first harvest, mixture-type plants performed better than monoculture-type plants in newly assembled monocultures of *P*. *vulgaris* and in mixtures of *G. mollugo* and *P. vulgaris* (Fig. 1a). However, in the two 267 mixtures with the small herbs V. chamaedrys and P. vulgaris and P. lanceolata and P.

268 *vulgaris*, monoculture-type plants performed better than mixture-type plants (see Fig. 1a).

269

# 270 3.2 Plant selection history and biodiversity effects

Overall, biodiversity effects were positive at both harvests (Fig. 2, Table 2).
Communities of naïve plants at the first harvest showed larger SEs and at the second harvest
showed smaller NEs and CEs than communities of selected plants.

274 At the first harvest, NEs, CEs and SEs were also larger for communities assembled 275 from monoculture-type plants than for communities assembled from mixture-type plants 276 (Fig. 2a, Table 2a). In contrast, at the second harvest NEs, CEs and SEs were non-277 significantly larger for communities assembled from mixture-type plants for most species 278 assemblages (Fig. 2b; Table 2b). As with the results obtained for community productivity, 279 the influence of selection history on biodiversity effects also depended on the specific 280 species combination in mixture assemblages (Table 2). NEs were larger for mixture-type 281 plants for the combinations of G. mollugo with either P. vulgaris or P. lanceolata at the first 282 harvest (Fig. A2 in Supplementary material; Table 2). At the second harvest, NEs and CEs 283 were generally more similar between selection histories across different combinations and 284 variation between the specific community compositions was mainly due to different SEs. 285 The much larger NE for mixture-type plants in the combination G. mollugo + P. vulgaris 286 was an exception (Fig. A2). For both harvests communities which included the legume L. 287 pratensis or the small herb P. lanceolata showed positive biodiversity effects (Fig. A2). Four 288 species combinations shifted between harvests from stronger CEs for monoculture-type 289 plants to stronger biodiversity effects for mixture-type plants (Fig. A2). The G. mollugo + P. 290 vulgaris combination showed a consistently larger CE for mixture-type plants. At the second harvest the different species combinations varied strongly in SEs, but not in CEs (Table 2).

292 SEs were often larger for mixture- than for monoculture-type plants (Fig. 2).

293

# 294 3.3 Plant selection history and within- and between-species trait variance

Whereas interspecific differences in plant height were marginally larger in mixturetype plants, interspecific differences in leaf thickness were larger in monoculture-type plants at the first harvest (Fig. 3). Intraspecific differences in SLA were larger for monoculture-type plants. Furthermore, pot-level SDs in monoculture or mixture assemblages were nonsignificantly larger for assemblages with monoculture- than with mixture-type plants (left two columns in Fig. 3).

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#### 302 *3.4 Relationship between biodiversity effects and trait variation*

303 At the first harvest, the NE was negatively correlated with the RD of plant height but 304 positively correlated with the RD of leaf thickness (Fig. 4). Selection history had a 305 significant effect on the relationship between SE and the RD of plant height but no or only 306 marginal effects on all other relationships. SEs were more negatively correlated with the RDs 307 of plant height for mixture- than for monoculture-type or naïve plants. In contrast, the RDs 308 of leaf thickness were positively correlated with NEs and CEs for both monoculture- and 309 mixture-type plants, but not for naïve plants (Fig. 4c). At the second harvest, NEs and SEs 310 were significantly negatively correlated with the RD of plant height (Fig. 5). CEs were not 311 influenced by interspecific variation in plant height.

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# 313 *3.5 Relationship between biodiversity effects and trait means*

Whereas the CE was negatively correlated with the CWM of SLA (Fig. 6b), the SE was positively correlated with the CWM of SLA (Fig. 6b, right panel). Consequently, the

316 NE, driven by the CE, decreased with increasing SLA. Leaf thickness had a marginally 317 significant effect on SE, but the directionality depended on selection history. Plant height did 318 not have a significant effect on the biodiversity effects at the first harvest. However, the 319 interaction between trait means and selection history was significant for the relationship 320 between the CWM of plant height and the SE at the first harvest. Selection history was not 321 significant for the relationship between biodiversity effects and CWMs for the other two 322 traits. At the second harvest, CWM of plant height had a significantly positive effect on NE, 323 CE and SE (Fig. 7), the biodiversity effects were therefore stronger for taller plants. 324 However, in contrast to the first harvest, at the second harvest no effect of selection history 325 on the relationship between the CWM of plant height and the SE was observed (Fig. 7).

326

#### 327 4. Discussion

328 *4.1 Influence of plant selection history on community productivity* 

329 Plant community productivity may be influenced by selection for increased niche 330 differentiation in plants grown for eight years in mixtures (mixture-type plants) compared 331 with plants grown in monocultures (monoculture-type plants, Zuppinger-Dingley et al., 332 2014). We hypothesized that 2-species mixtures comprised of mixture-type plants should 333 have greater community productivity than the same mixtures comprised of monoculture-type 334 plants. Conversely, we expected monocultures with monoculture-type plants to have greater 335 community productivity than the same monocultures with mixture-type plants. For naïve 336 plants, we expected intermediate community productivity in both monocultures and 337 mixtures.

Our results provide mixed evidence for these hypotheses, in part depending on the particular species and species combinations. Plant assemblages consisting of naïve plants, without a selection history in the Jena Experiment, generally produced the lowest community biomass, especially in 2-species mixtures, in the pots in the glasshouse in Zurich. Evolutionary processes in the field plots likely led to the increased performance of selected plants, because these plants were grown for a longer time without re-sowing. In contrast, the naïve plants were re-sown every year in the commercial propagation cultures, thereby "resetting" any local adaptation with every generation.

346 Within the selected plants, mixture-type plants produced higher community biomass 347 than monoculture-type plants in 2-species mixtures. But mixture-type plants also produced 348 more biomass than monoculture-type plants when grown in monoculture, which reduced 349 biodiversity effects. The generally lower performance of monoculture-type plants may have 350 been due to selection for increased defense against pathogens that are known to accumulate 351 in monocultures (Schnitzer et al., 2011). Increased resource allocation to defense 352 mechanisms would result in reduced allocation to growth (Coley et al., 1985; Herms and 353 Mattson, 1992). However, during the 24 weeks of growth in the experiment, species-specific 354 pathogens may not have been present, removing the need for increased defense. Such 355 species-specific pathogens might have needed more time to accumulate and to render 356 increased pathogen resistance advantageous.

357 Selection-history effects varied strongly among species in monocultures and among 358 species compositions in 2-species mixtures. These findings emphasized that such studies 359 should include multiple species but at the same time sufficient replication in monoculture 360 and their combinations in mixture. High replication can more easily be achieved in 361 experiments with one focal species (e.g. Kleynhans et al., 2016; Rottstock et al., 2017), but 362 extrapolating results from such experiments might under- or overestimate overall effects of 363 selection on the response of plants to different biotic conditions. We used five focal species 364 and observed strong differences regarding their selection response to community diversity.

365

#### 366 4.2 Influence of plant selection history on biodiversity effects

Net biodiversity effects can be partitioned into CEs and SEs. When CEs drive overyielding, most species contribute similarly to greater community productivity in mixtures, presumably due to niche differentiation among them. Conversely, SEs are large when few dominant species are driving positive diversity–productivity relationships, because they benefit from growing in mixtures (Loreau and Hector 2001).

372 Naïve plants exhibited weak biodiversity effects, confirming findings from a field 373 experiment, where biodiversity effects were weaker for assemblages of naïve plants, 374 especially at low diversity levels (van Moorsel et al., 2017). Such naïve plants, in contrast to 375 plants with a common selection history, did not experience continued selection in field plots 376 without re-sowing and did not previously experience interspecific competition. In contrast to 377 our expectations, not only NEs and SEs but also CEs were larger for monoculture- than for 378 mixture-type plant communities at the first harvest. However, the lower CEs of mixture-type 379 communities could be attributed to a higher performance of mixture-type plants in 380 monoculture assemblages and not to a lower performance in mixed assemblages. At second 381 harvest, NEs, CEs and SEs were similar for mixture- and monoculture-type communities. 382 Nevertheless, at least in four 2-species combinations — L. pratensis + V. chamaedrys, G. 383 mollugo + V. chamaedrys, V. chamaedrys + P. vulgaris and P. lanceolata + P. vulgaris — 384 the directionality changed, i.e. CEs at the second harvest were larger for mixture- than for 385 monoculture-type communities. Over longer timespans, CEs often increase and SEs often 386 decrease (Fargione et al., 2007; Isbell et al., 2009; Marquard et al., 2009; Montès et al., 2008; 387 van Ruijven and Berendse, 2005). Possibly this would have occurred in our experiment if we 388 had continued beyond the 24-weeks of the study.

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#### 391 *4.3 Influence of plant selection history on trait variation*

392 Because community-level trait variation can reflect niche differentiation (Roscher et 393 al., 2015; Violle et al., 2012), we measured intra- and interspecific trait variation among 394 individual plants in all assemblages. We expected larger interspecific trait variation for 395 mixture-type plants undergoing possible selection for increased complementarity during 396 twelve years in the experimental field plots. Conversely, we expected stronger within-species 397 trait variation in monoculture-type plants with twelve years of strong intraspecific 398 competition in the experimental field plots. However, monoculture-type plants tended to 399 show higher intra and inter-specific trait variation (see Fig. 3). The relative extent of 400 intraspecific trait variation may depend on species richness (Hulshof et al., 2013; Lamanna et 401 al., 2014; Le Bagousse-Pinguet et al., 2014; Siefert et al., 2015) and in monocultures a large 402 intraspecific variation is advantageous for a more efficient use of resources. Thus, the trend 403 for increased trait variation in monoculture-type plants is consistent with potential selection 404 for intraspecific niche widening by within-species character displacement during prolonged 405 growth in monocultures.

406 However, less interspecific trait variation in mixture- compared with monoculture-407 type plants was in accordance with the lower CEs for mixture-type plants. These findings 408 contrast with an earlier study in which larger CEs were observed for mixture- than for 409 monoculture type plants and where mixture-type plants showed increased interspecific trait 410 variation (Zuppinger et al., 2014). This earlier study included more species that were 411 functionally different from each other, namely grasses, legumes, small herbs and tall herbs, 412 which could potentially explain the contrasting results. More similar species, such as those 413 used in the present study, may not be able to further increase trait differences in such a short 414 time frame (Allan et al., 2013). Such species may have evolved "parallel" character 415 displacement in response to species of the other functional groups also present in the 416 mixtures in which they were selected in the Jena Experiment.

417

# 418 4.4 Influence of trait variation and community-weighted means on biodiversity effects

419 Selection for niche differentiation (Zuppinger-Dingley et al., 2014) could explain the 420 increase of biodiversity effects over time in field experiments (Cardinale et al., 2007, Reich 421 et al., 2012). Not all trait variation, however, corresponds to niche differentiation (Turcotte 422 and Levine 2016). In particular, traits related to light availability may behave differently 423 because of the asymmetric nature of competition for light, i.e. being tall is generally better 424 than being small. Thus, variation in plant height could be expected to decrease when species 425 are grown in mixtures rather than monocultures (Roscher et al., 2015; Vermeulen et al., 426 2008). In the present study, the relationship between functional traits in 2-species mixtures 427 and biodiversity effects did not consistently differ between plants selected in monocultures 428 vs. mixtures and this was consistent with the absence of increased CEs and between-species 429 trait variations in mixture-type plants. Nevertheless, we did test for such relationships — 430 independent of selection treatments — between functional traits and biodiversity effects in 431 our 2-species mixtures. Specifically, we tested whether relative trait differences (RD) were 432 positively correlated with CEs and community-weighted trait means (CWMs) were 433 positively correlated with SEs.

RDs in plant height were negatively rather than positively correlated with CEs and consequently NEs (see Fig. 4a, 5). This discrepancy between our expectation and observation suggests that RDs in plant height may reflect competitive hierarchies rather than complementary of plants with respect to light use, as discussed above with regard to the asymmetry of light competition. At the second harvest, CWMs of plant height had a positive impact on all biodiversity effects (Fig. 7), i.e. not only as has been previously observed on SEs (Cadotte 2017; Roscher et al., 2015; Vermeulen et al., 2008). 441 Functional diversity in SLA within a community should increase complementary light 442 use (Roscher et al., 2011). Leaf thickness is inherently related to SLA (White and Montes-R, 443 2005) and might act similarly to SLA. Here, RDs in leaf thickness, but not RDs in SLA, 444 were positively correlated with all biodiversity effects, especially for mixture-type plants. 445 Trait plasticity in leaf thickness was therefore advantageous for species growing in mixtures. 446 However, SEs increased as much as CEs, contrary to our expectation that positive 447 correlations between trait differences may mainly involve CEs. Furthermore, CWMs of SLA 448 had a positive effect on SEs, but also a negative effect on CEs, adding up to a negative effect 449 on NEs, suggesting that a smaller leaf area per unit mass for species growing in mixtures has 450 a positive effect on productivity.

451

#### 452 **5.** Conclusions

453 Here, we demonstrated that community diversity had the selective potential to alter 454 species performances, which may in part explain the strengthening of biodiversity-455 ecosystem functioning relationship observed in field experiments (e.g. Reich et al., 2012). 456 Selection in a biodiversity experiment increased community productivity in newly assembled 457 test assemblages compared to assemblages composed of naïve plants without such selection 458 history. Moreover, previous selection in mixtures increased community productivity in 459 newly assembled mixtures and monocultures compared with previous selection in 460 monocultures. These findings imply that co-evolutionary processes occurred in the 12-year 461 selection period in the experimental plots of the biodiversity experiment and involving at 462 least two sexual reproduction cycles. Such rapid evolutionary processes in grassland plant 463 communities have implications for conservation strategies. Thus, it may not be sufficient to 464 only conserve species in isolation but rather in communities or populations of species with 465 co-evolved interactions.

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#### 478 Supplementary data

479 Supplementary data associated with this article can be found in the online version.

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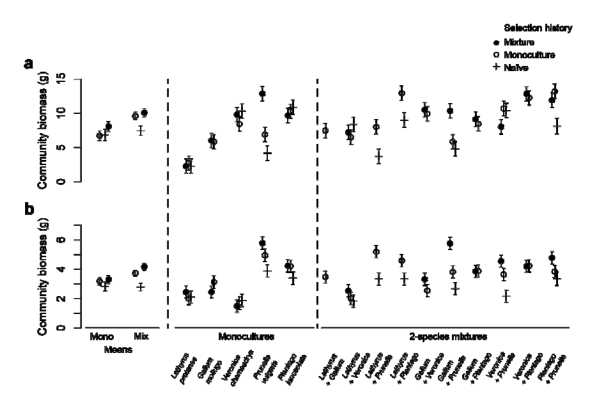
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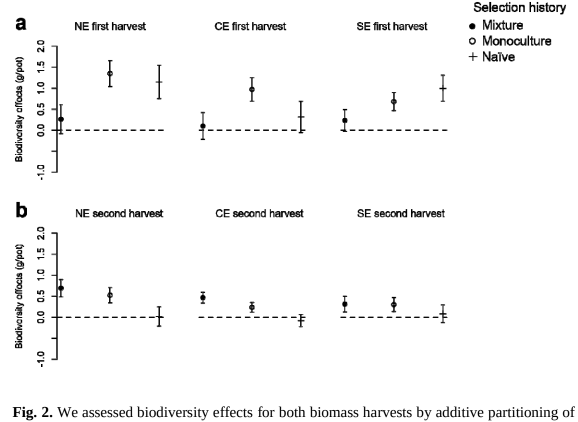
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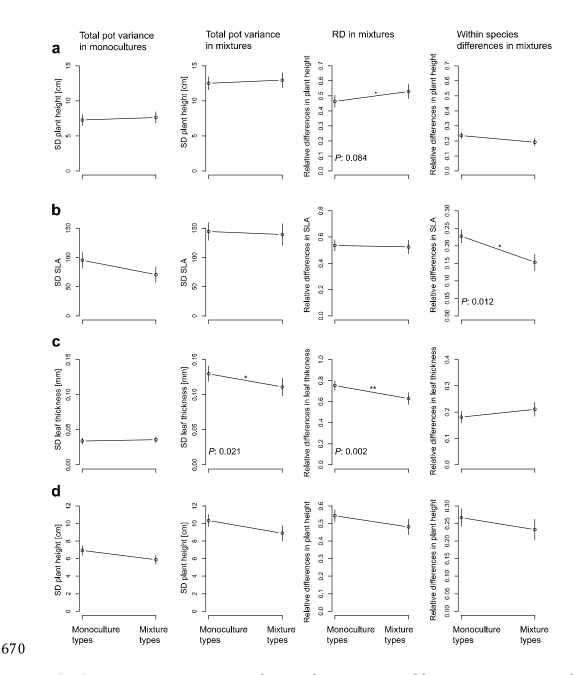
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Fig. 1. Mean community biomass for monocultures and 2-species mixtures. Shown are means and standard errors from a linear mixed-effects model with selection history, species combination and the interaction between selection history and species assembly as fixedeffects terms and table (including the block) as random-effects term. **a**, first harvest. **b**, second harvest.

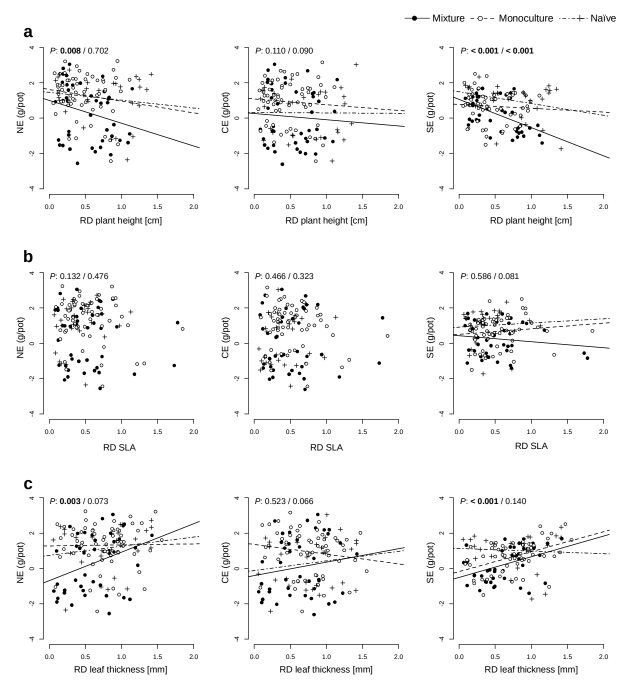


**Fig. 2.** We assessed biodiversity effects for both biomass harvests by additive partitioning of the net effect (**a**, NE) into complementarity effect (**b**, CE) and sampling effect (**c**, SE) for plants with different selection histories (naïve, monoculture, mixture). Shown are means and standard errors from a linear mixed-effects model, with selection history as fixed-effects term and species assembly, the interaction between selection history and species assembly and table (including block) as random-effects terms.

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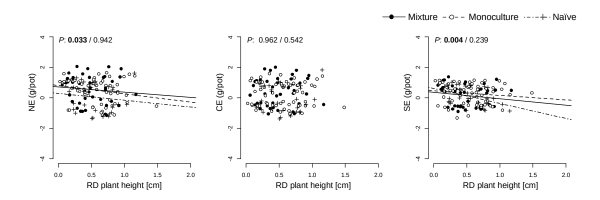
**Fig. 3.** Trait variance in monoculture and mixture assemblages in response to selection history (monoculture- vs. mixture-type plants). **a**) plant height at first harvest, **b**) SLA at the first harvest, **c**) leaf thickness at the first harvest, **d**) plant height at second harvest. Shown are means and standard errors from a mixed-effects model with selection history, species assembly and the two-way interaction of these as fixed-effects terms and table (including block) as random term. Significant and marginally significant *P*-values are indicated in the respective plot.



**Fig. 4**. Biodiversity effects at the first harvest in response to relative differences between species (RDs) for three traits: **a**, plant height (in cm), **b**, specific leaf area (SLA) and **c**, leaf thickness (in mm). Indicated *P*-values refer to ANOVA results for fixed-effects terms from a mixed-effects model with RD, species assembly, selection history and interactions of these

- 683 as fixed-effects terms and table (including block) as random-effects term: RD / interaction
- 684 RD × selection history (naïve plants vs. mixture types vs. monoculture types). Regression
- 685 lines are plotted in cases for which at least one *P*-value was significant. Left column: NE,
- 686 middle column: CE, right column: SE.

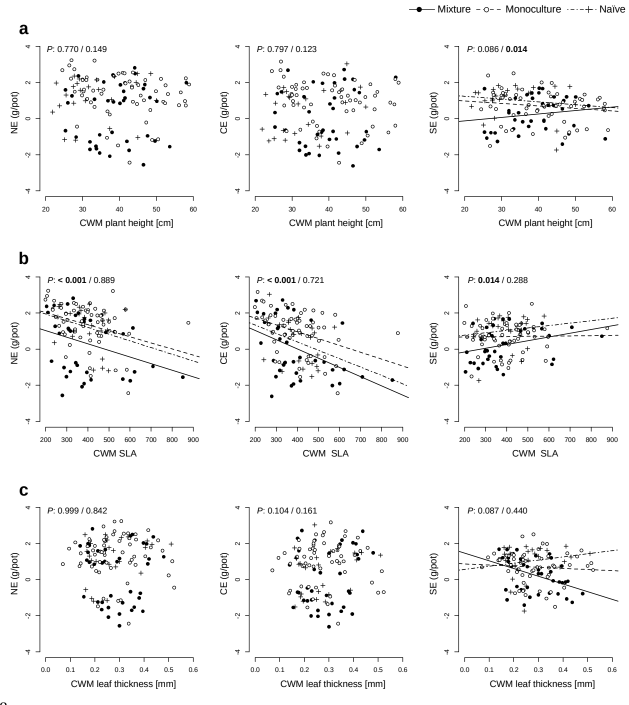
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**Fig. 5.** Biodiversity effects at the second harvest in response to relative differences between species for plant height (in cm). Indicated *P*-values refer to ANOVA results for fixed-effects terms from a mixed-effects model with RD, species assembly, selection history and interactions of these as fixed-effects terms and table (including block) as random-effects term: RD / interaction RD × selection history (naïve plants vs. mixture types vs. monoculture types). Regression lines are plotted in cases for which at least one *P*-value was significant. Left column: NE, middle column: CE, right column: SE.

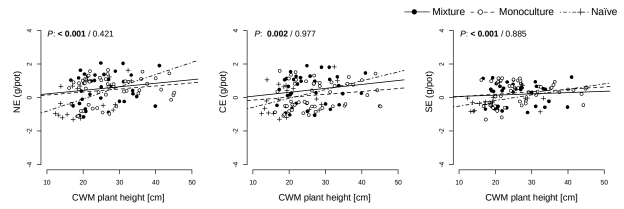
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**FIG. 6.** Biodiversity effects at the first harvest in response to the community-weighted mean (CWM) of three traits: **a**, plant height (in cm), **b**, specific leaf area (SLA) and **c**, leaf thickness (in mm). Indicated *P*-values refer to ANOVA results for fixed-effects terms from a mixed-effects model with CWM, species assembly, selection history and interactions of these as fixed-effects terms and table (including block) as random-effects term: CWM /

- interaction CWM × selection history (naïve plants vs. mixture types vs. monoculture types).
- 706 Regression lines are plotted in cases for which at least one *P*-value was significant. Left
- 707 column: NE, middle column: CE, right column: SE.



**Fig. 7.** Biodiversity effects at the second harvest in response to the community-weighted mean (CWM) of plant height (in cm). Indicated *P*-values refer to ANOVA results for fixedeffects terms from a mixed-effects model with CWM, species assembly, selection history and interactions of these as fixed-effects terms and table (including block) as random-effects term: CWM / interaction CWM × selection history (naïve plants vs. mixture types vs. monoculture types). Regression lines are plotted in cases for which at least one *P*-value was significant. Left column: NE, middle column: CE, right column: SE.

# **Table 1.** Results of mixed-effects ANOVA for the aboveground biomass of the test assemblages at first harvest after 12 weeks of growth (a) and

718 at second harvest after 24 weeks of growth (b).

|                                    |  |     | a) Harvest 1 |       |         | b) Harvest 2 |       |         |
|------------------------------------|--|-----|--------------|-------|---------|--------------|-------|---------|
| Source of variation                |  | nDf | dDF          | F     | Р       | dDF          | F     | Р       |
| Species assembly:                  |  |     |              |       |         |              |       |         |
|                                    | Monoculture vs. mixture                                      | 1   | 173.3        | 29.09 | < 0.001 | 174          | 10.78 | 0.001   |
|                                    | Monoculture identity or species combination of mixture       | 13  | 171.2        | 16.53 | < 0.001 | 171.8        | 15.47 | < 0.001 |
| Selection history:                 |  |     |              |       |         |              |       |         |
|                                    | Naïve vs. mono or mix types                                  | 1   | 173          | 16.63 | < 0.001 | 173.7        | 42.72 | < 0.001 |
|                                    | Mono vs. mix types   | 1   | 169.6        | 1.78  | 0.184   | 170.1        | 5.71  | 0.018   |
| Assembly $\times$ history:         |  |     |              |       |         |              |       |         |
|                                    | Monoculture vs. mixture $\times$ naïve vs. mono or mix types | 1   | 168.4        | 1.72  | 0.191   | 168.8        | 8.56  | 0.004   |
|                                    | Monoculture vs. mixture $\times$ Mono or mix types           | 1   | 172.2        | 1.69  | 0.195   | 172.9        | 3.52  | 0.062   |
| Species assembly $\times$ history: |  |     |              |       |         |              |       |         |
|                                    | Species assembly $\times$ naïve vs. mono or mix types        | 8   | 171.7        | 5.35  | < 0.001 | 172.3        | 2.15  | 0.033   |
|                                    | Species assembly $\times$ mono types vs. mix types           | 10  | 172.3        | 2.91  | 0.002   | 172.9        | 1.23  | 0.275   |
|                                    |  | _   |              |       |         |              | _     |         |

nDF = numerator degrees of freedom, dDF = denominator degrees of freedom, F = variance ratio, P = probability of type-I error. Variance components (Var) and associated standard errors (SE) for the random effects are provided together with the number of replicates.

720

#### 722 Table 2. Results of mixed-effects ANOVA for biodiversity effects of the test assemblages at

the first harvest after 12 weeks of growth (a) and at the second harvest after 24 weeks of

724 growth (**b**).

|   |     | a) NE | Harvest | 1       | b) NE Harvest 2 |       |         |
|---|-----|-------|---------|---------|-----------------|-------|---------|
| Source of variation                                   | nDf | dDF   | F       | Р       | dDF             | F     | Р       |
| Overall mean  | 1   | 15.9  | 26.67   | < 0.001 | 15.1            | 14.35 | 0.002   |
| Naïve vs. mono or mix types                           | 1   | 95.3  | 0.806   | 0.372   | 96.1            | 11.54 | < 0.001 |
| Mono types vs. mix types                              | 1   | 93.9  | 21.01   | < 0.001 | 93.6            | 0.026 | 0.872   |
| Species assembly                                      | 9   | 96.7  | 2.646   | 0.009   | 97.7            | 4.837 | < 0.001 |
| Species assembly $\times$ naïve vs. mono or mix types | 4   | 97.5  | 4.459   | 0.002   | 98.9            | 1.463 | 0.219   |
| Species assembly $\times$ mono types vs. mix types    | 6   | 98    | 4.095   | 0.001   | 99              | 1.518 | 0.18    |

|   |     | a) Cl | E harvest | 1       | b) CE Harvest 2 |       |       |  |
|---|-----|-------|-----------|---------|-----------------|-------|-------|--|
| Source of variation                                   | nDf | dDF   | F         | Р       | dDF             | F     | Р     |  |
| Overall mean  | 1   | 15.8  | 8.214     | 0.011   | 14.5            | 4.108 | 0.061 |  |
| Naïve vs. mono or mix types                           | 1   | 95.9  | 1.427     | 0.235   | 96.6            | 5.668 | 0.019 |  |
| Mono types vs. mix types                              | 1   | 94.4  | 14.2      | < 0.001 | 93.8            | 1.524 | 0.22  |  |
| Species assembly                                      | 9   | 97.4  | 2.534     | 0.012   | 98.4            | 1.121 | 0.356 |  |
| Species assembly $\times$ naïve vs. mono or mix types | 4   | 98.3  | 1.835     | 0.128   | 99.7            | 0.584 | 0.675 |  |
| Species assembly $\times$ mono types vs. mix types    | 6   | 98.8  | 2.53      | 0.025   | 99.8            | 0.468 | 0.831 |  |

|   |     | a) SE harvest 1 |       |         |       | b) SE Harvest 2 |         |  |
|---|-----|-----------------|-------|---------|-------|-----------------|---------|--|
| Source of variation                                   | nDf | dDF             | F     | Р       | dDF   | F               | Р       |  |
| Overall mean  | 1   | 14.2            | 97.07 | < 0.001 | 15.1  | 11.66           | 0.004   |  |
| Naïve vs. mono or mix types                           | 1   | 104.2           | 12.66 | 0.001   | 98.8  | 2.224           | 0.139   |  |
| Mono types vs. mix types                              | 1   | 101.2           | 10.28 | 0.002   | 95.7  | 2.37            | 0.127   |  |
| Species assembly                                      | 9   | 105.5           | 5.793 | < 0.001 | 100.8 | 11.53           | < 0.001 |  |
| Species assembly $\times$ naïve vs. mono or mix types | 4   | 105.9           | 10.08 | < 0.001 | 102   | 3.517           | 0.01    |  |
| Species assembly $\times$ mono types vs. mix types    | 6   | 105.9           | 2.865 | 0.013   | 101.9 | 2.541           | 0.025   |  |

nDF = numerator degrees of freedom, dDF = denominator degrees of freedom, F = variance ratio, P = probability of type-I error. Variance components (Var) and associated standard errors (SE) for the random effects are provided together with the number of replicates. 725