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PLANT COMPETITION IN RELATION TO NEIGHBOR BIOMASS: AN INTERCONTINENTAL STUDY WITH *POA PRATENSIS*¹

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Abstract. A standardized neighbor removal experiment was conducted in 12 plant communities located on three continents to test the null hypothesis that competition intensity (CI) was independent of the amount of plant biomass present. Six plots were chosen in each community to cover the range of local variation in plant biomass. In each plot the relative growth rate (RGR) of transplanted *Poa pratensis* (Poaceae) seedlings was compared in the presence and absence of neighbors. Neighbors were removed experimentally using

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herbicide. Removing neighbors increased RGR of transplants significantly in most plots. CI increased with an increase in the amount of neighbor biomass present in one community where the range of neighbor biomass was greater than in any other community. In contrast, CI did not change significantly with an increase in neighbor biomass in other communities where the range of neighbor biomass was smaller. For the communities combined, CI was not related to neighbor biomass in a consistent fashion. These results indicate that competition may reduce growth over a wide range of habitat productivity, but the relationship between CI and neighbor biomass may differ among communities.

Key words: Australia; Europe; habitat productivity; North America; plant competition; *Poa pratensis*; removal experiment.

INTRODUCTION

Current theories of plant community organization differ in their prediction about the relationship between competition intensity (CI) and habitat productivity. Some theories (Grime 1979, Keddy 1990) predict that competition intensifies as productivity increases because of a corresponding increase in plant biomass. Other theories (Tilman 1988, Taylor et al. 1990) predict that competition does not intensify with an increase in habitat productivity for two different reasons. First, competition for light may intensify with an increase in habitat productivity but competition for soil nutrients and water may be less intense in productive habitats than in unproductive habitats; consequently, total competition should remain relatively constant along a productivity gradient (Newman 1973, Grubb 1985, Tilman 1988). Second, competition intensity depends on the ratio of resource supply to demand and this ratio may be unrelated to habitat productivity (Taylor et al. 1990).

A number of neighbor removal experiments have been conducted to test these conflicting predictions. In each experiment the response of a target plant to removal of its neighbors was compared at several positions on a productivity gradient. Experimental results have been inconsistent. In some experiments (e.g., del Moral 1983, Gurevitch 1986, Wilson and Keddy 1986, Reader and Best 1989), the performance of the target plant improved most where habitat productivity was greatest. In other experiments (e.g., Fowler 1990, Wilson and Shay 1990, Wilson and Tilman 1991, 1993), the performance of the target plant improved equally at all levels of productivity.

These experimental results may be inconsistent because CI was measured over different ranges of habitat productivity, using different target species, different experimental designs, and different methods to calculate CI. The chance of detecting a relationship between CI and habitat productivity may have been greater where a wider range of neighbor biomass values was considered. Unfortunately, it is difficult to check this possibility because studies involved different target species and interspecific variation in competitive ability is a potentially confounding factor. In the present study, we used the same target species in a standardized neighbor removal experiment, conducted in 12 plant communities on three continents, to test the null hypothesis that CI is independent of neighbor biomass. The null

hypothesis was tested for each community and for all communities combined. The target species used for the experiment was the perennial grass *Poa pratensis* L., henceforth called *Poa*. *Poa* was chosen for two reasons. First, it is widely naturalized and therefore appropriate for use in an intercontinental experiment. Second, its response to neighbor removal did not change significantly across local productivity gradients in previous studies (DiTommaso and Aarssen 1991, Wilson and Tilman 1991). Whether response of such a species to neighbor removal is independent of neighbor biomass over a wide range of habitat productivity also needs to be tested.

The theories mentioned above do not state explicitly how CI should be calculated. Consequently, CI has been equated with the absolute reduction in plant performance due to neighbors (e.g., plant growth rate without neighbors minus plant growth rate with neighbors) or with the relative reduction in plant performance due to neighbors (e.g., difference in plant growth rate without neighbors and with neighbors, divided by plant growth without neighbors) (Keddy 1989, Campbell and Grime 1992, Grace 1993, Turkington et al. 1993). A potential limitation of using the absolute index of CI in this case is that index values may increase with habitat productivity not only because competition intensifies as productivity increases (i.e., plant performance declines where neighbors are present, while performance is constant with neighbors absent) but also because there may be fewer environmental constraints on plant performance as productivity increases and plant performance may improve where neighbors are absent while performance is constant with neighbors present. The relative index of CI allows for possible interhabitat differences in plant performance where neighbors are absent by expressing the absolute reduction in plant performance due to neighbors relative to plant performance with neighbors absent. However, a potential limitation of the relative index of CI is that it may obscure differences in absolute reduction in plant performance due to neighbors. In the present study, both absolute and relative indices of CI were calculated.

METHODS

Study sites

The study was conducted at 12 locations (Table 1). At each location, a site was chosen that was appropriate

TABLE 1. Aboveground biomass of neighbors and dominant species at study sites, ordered by increasing neighbor biomass. $n = 6$ plots.

Location	Range of neighbor biomass (g/m ²)	Species	
		Four most abundant	(Mean Daubenmire score)†
1. Guelph, Canada (43°33' N, 80°10' W)	26–199	<i>Hieracium floribundum</i> <i>Poa pratensis</i> <i>Poa compressa</i> <i>Viccia cracca</i>	(3) (3) (2) (2)
2. Almonte, Canada (45°15' N, 76°15' W)	61–246	<i>Carex crawei</i> <i>Sporobolus heterolepis</i> <i>Danthonia spicata</i> <i>Sporobolus vaginiflorus</i>	(3) (2) (1) (1)
3. Canberra, Australia (35°17' S, 149°11' E)	121–234	<i>Themeda australis</i> <i>Stipa falcata</i> <i>Danthonia carphoides</i> <i>Taraxacum officinale</i>	(3) (2) (2) (1)
4. Morgantown, USA (39°39' N, 79°59' W)	155–194	<i>Solidago</i> spp. <i>Danthonia spicata</i> <i>Vernonia novaboracensis</i> <i>Impatiens pallida</i>	(–) (–) (–) (–)
5. Vancouver, Canada (49°15' N, 123°14' W)	174–302	<i>Polygonum persicaria</i> <i>Trifolium repens</i> <i>Stellaria</i> spp. <i>Gnaphalium uliginosum</i>	(1) (1) (1) (1)
6. Umeå, Sweden (64°12' N, 17°34' E)	158–384	<i>Geranium sylvaticum</i> <i>Agrostis capillaris</i> <i>Cirsium helenoides</i> <i>Alchemilla</i> sp.	(3) (2) (2) (1)
7. Long Island, USA (40°55' N, 73°8' W)	163–436	<i>Solidago altissima</i> <i>Solidago rugosa</i> <i>Poa annua</i> <i>Viccia cracca</i>	(–) (–) (–) (–)
8. Sawyerville, Canada (45°25' N, 71°30' W)	237–557	<i>Festuca rubra</i> <i>Phalaris arundinacea</i> <i>Agrostis stolonifera</i> <i>Cerastium arvense</i>	(3) (3) (1) (1)
9. Groningen, The Netherlands (53°5' N, 6°40' E)	81–648	<i>Agrostis stolonifera</i> <i>Holcus lanatus</i> <i>Plantago lanceolata</i> <i>Equisetum arvense</i>	(2) (1) (1) (1)
10. Baton Rouge, USA (30°25' N, 91°10' W)	‡	<i>Andropogon glomeratus</i> <i>Andropogon virginicus</i> <i>Paspalum notatum</i> <i>Sporobolus indicus</i>	(3) (2) (1) (2)
11. Minneapolis, USA (45°24' N, 93°12' W)	‡	<i>Hedeoma hispida</i> <i>Aristida basiramea</i> <i>Ambrosia coronopifolia</i> <i>Panicum oligosanthes</i>	(1) (1) (1) (1)
12. Sydney, Australia (33°56' S, 151°12' E)	‡	<i>Cynodon dactylon</i> <i>Pennisetum clandestinum</i> <i>Chloris gayana</i> <i>Stenotaphrum secundatum</i>	(3) (1) (1) (1)

† 1 = 0–5%, 2 = 6–25%, 3 = 26–50% cover, – = cover not recorded.

‡ Neighbor biomass not recorded.

for *Poa* (e.g., grassland or old field) and where there was a productivity gradient due to within-site variation in factors such as soil depth, soil fertility, topography, or land use practices.

Cover of naturally occurring vascular plants at each site was recorded in 20 0.5 × 1 m plots using the six class Daubenmire scale (Mueller-Dombois and Ellenberg 1974). Plots were chosen subjectively to account

for variation in productivity (i.e., amount of vegetation present) at each site. A mean cover score was calculated for each species at each site.

Experimental design and procedure

At each site, an additional six plots, each ≈ 13 m², were chosen subjectively, based on the amount of vegetation present, to include the range of within-site vari-

ation in productivity. Plots were at least 30 m apart. Within each plot 10 0.5 m diameter subplots (1 m apart) were chosen subjectively; 5 subplots were assigned randomly as controls (i.e., vegetation left intact) and 5 were treated in early spring with a contact herbicide (glyphosate [N-phosphonomethyl glycine], trade name Roundup) to remove vegetation. The herbicide was contained during application by enclosing the treated subplot in a 0.5 m diameter \times 1 m tall metal cylinder that was held firmly to the ground. Possible translocation of herbicide was prevented by severing roots around the perimeter of each treated subplot to a depth of 15 cm with a shovel. Root severing was repeated in treated subplots at 3-wk intervals throughout the experiment to minimize invasion of roots of neighboring plants. Treated subplots were kept free of shoots of neighboring plants by hand-weeding. To exclude large herbivores such as rabbits, wire screening (1.25 cm mesh, 33 cm tall) was placed around the perimeter of all subplots.

Response to neighbor removal

Seeds from a single seedlot of *Poa pratensis* var. Nassau were used for the experiment. Seeds were collected from plants grown in field plots at 1 of the 12 locations (i.e., Guelph, Canada). At each location, seeds were sown in 2.5 \times 2.5 \times 14 cm deep pots (Roottrainers, Spencer-Lemaire Industries, Alberta, Canada) that were filled with a 1:6 mixture of local soil: peat. After seeds germinated, seedlings were thinned to one per pot and they were grown in a greenhouse at each location to achieve a standard size (shoot mass of 0.1 g) for transplanting into field plots. Seedling shoot mass was estimated using a regression equation in which shoot mass was the dependent variable and total leaf length and (or) leaf number were independent variables. At each location, the regression equation was constructed from data collected for 20 plants grown in the greenhouse together with transplants.

One seedling was transplanted into the center of each of the 60 subplots per location (10 subplots \times 6 plots). During the first 2 wk after transplanting, seedlings were shaded by a 30 \times 40 cm piece of 70% shade cloth and were watered to minimize mortality. Dead seedlings were replaced during the first 3 wk after transplanting.

After 10–12 wk, shoots of transplants and total aboveground biomass of neighbors in control plots were harvested, dried, and weighed. Belowground tissues were not harvested because of the practical difficulty of collecting small roots.

The relative growth rate (RGR) of each transplant was calculated as

$$\text{RGR} = \frac{\ln(M_2) - \ln(M_1)}{t_2 - t_1},$$

where M_2 is shoot mass at harvest, M_1 is estimated shoot mass at transplanting, and $t_2 - t_1$ is the number of days between harvest date (t_2) and transplanting date

(t_1). Mean RGR was calculated for treated and control subplots in each plot. For each site, the statistical significance of a difference between values of mean RGR for treated and control subplots was tested using Student's t test. The total number of transplants surviving to the end of the experiment was also compared for treated and control subplots at each site. The statistical significance of a difference in transplant survival was tested using a G test (Sokal and Rohlf 1981).

Competition intensity

Mean values of RGR were used to calculate both absolute competition intensity (CI_a) and relative competition intensity (CI_r). CI_a was calculated as $\text{RGR}_t - \text{RGR}_c$ and CI_r was calculated as $(\text{RGR}_t - \text{RGR}_c)/\text{RGR}_c$, where RGR_t and RGR_c are the mean relative growth rates of transplants in treated subplots (i.e., neighbors removed) and control subplots (i.e., neighbors left intact), respectively.

For each index, CI values were first analyzed separately for each site to test the null hypothesis that CI was independent of neighbor biomass. Least squares linear regression was used to test the null hypothesis (i.e., slope of regression was not significantly different from zero [$P > 0.05$]). CI values for all sites were then combined to test the null hypothesis that CI was independent of neighbor biomass over a wider range of neighbor biomass values. Generalized additive modelling (Hastie and Tibshirani 1990) was used to test for a statistically significant relationship between CI and neighbor biomass for all sites combined. This technique was chosen because it can be used for both linear modelling and nonlinear modelling. Our model contained a continuous predictor (neighbor biomass) as well as a categorical factor (site). If the relationship between the continuous predictor and the response is assumed to be linear, then the model is an analysis of covariance (ANCOVA), if nonlinear, a generalized additive model (GAM). In both cases, for the model to be interpretable, one has to assume a lack of interaction between effects of the continuous predictor and the categorical factor. This assumption was tested and verified.

In GAM, the most appropriate functional form of the relationship between the dependent variable and each independent variable is derived from the data using scatterplot smoothing (see Hastie and Tibshirani 1990 for details). The form of the model used here was

$$Y = a + b_s + cf(\text{neighbor biomass}) + \text{error},$$

where a is the general intercept, b_s is the intercept for site s , c is a constant, and f is a smoothing function for neighbor biomass. The constants a , b_s , and c are estimated from the data. If f is taken to be the identity function then the model reduces to a linear one (ANCOVA).

The statistical significance of a particular independent variable, neighbor biomass in this case, is assessed

TABLE 2. Survival of *Poa* transplants in $n = 6$ experimental plots.

Location	Number surviving ($n = 30$)		Number of plots with low* survival for sub- plots with neighbors removed
	Neighbors left intact	Neighbors removed	
1. Guelph	21	30	0
2. Almonte	24	22	3
3. Canberra	25	28	0
4. Morgantown	15	15	3
5. Vancouver	25	24	2
6. Umeå	30	30	0
7. Long Island	26	24	1
8. Sawyerville	30	29	0
9. Groningen	24	25	1
10. Baton Rouge	6	1	6
11. Minneapolis	9	7	6
12. Sydney	7	7	6

* Less than four of five transplants per subplot survived.

using the reduction in residual deviance (D) attributable to a model with the variable included, as compared to a model excluding the variable. For our data, with an identity function and assumed Gaussian remainders, D reduces to the remainder sum of squares (Hastie and Tibshirani 1990). A reduction in D is tested for statistical significance by comparing the reduction in D to a chi-square distribution with the degrees of freedom (df) for the simple model minus df for the more complex model. In this case, the simple model had site as the independent variable and the more complex model had both site and neighbor biomass as independent variables. In GAM, the degree of smoothing applied to fit a possibly nonlinear relationship can be controlled by setting the degrees of freedom for the continuous predictor; df = 1 specifies a linear fit, df = 5 specifies a very nonlinear fit. Whether the nonlinear fit is a significant improvement over the linear fit is tested in the usual way by comparing the two models. In this case, the statistical significance (i.e., P values) of the variables neighbor biomass and site did not change appreciably when ANCOVA rather than GAM was used so only GAM results are reported here.

The proportion of variation (R^2) accounted for by a particular GAM is calculated as the difference between D for a model containing only a constant term (null model) and D for the model under consideration, divided by D for the null model.

Our analysis standardized for potential site effects on CI by modelling CI as the sum of a site factor plus a (possibly nonlinear) function of neighbor biomass. One index of competition intensity (CI_c) also standardized for potential site effects by expressing CI as a ratio. These two standardization procedures are complementary so GAM was used to test the null hypothesis for CI_c , as well as for CI_a .

RESULTS

Neighbor biomass and species composition

Neighbor biomass and species composition both varied considerably among sites (Table 1). Mean neighbor biomass per plot ranged from a low of 26–199 g/m² at Guelph to a high of 81–648 g/m² at Groningen. Vegetation was dominated by graminoids and forbs at each site but there was little overlap in dominant species among sites (Table 1). The target species *Poa* occurred naturally at four sites (Guelph, Vancouver, Umeå, and Groningen).

Response to neighbor removal

Transplant survival in subplots with neighbors left intact ranged from 6 to 30 plants of the 30 plants added per site (Table 2). Removing neighbors increased transplant survival significantly ($P < 0.05$, G test) only at one site (Guelph).

Survival was low even in the absence of neighbors at some sites (Table 2). Drought and herbivores that were not excluded by the wire screening (e.g., insects, rodents) killed almost all transplants at three sites (Baton Rouge, Minneapolis, and Sydney). The same factors contributed to low survival in one to three of the six plots at each of five other sites (Almonte, Morgantown, Vancouver, Long Island, and Groningen). In total, 28 of the 72 plots, i.e., 6 plots \times 12 sites, had low survival of transplants in the absence of neighbors. Factors in addition to competition from neighbors probably limited transplant performance in these 28 plots so they were excluded from all subsequent analyses.

In the remaining 44 plots, mean RGR ranged from -0.016 to 0.024 g·g⁻¹·d⁻¹ in subplots with neighbors left intact (Fig. 1). RGR was negative in 17 of 44 subplots because transplants lost mass during the experiment. In subplots with neighbors removed, mean RGR was positive in all cases and ranged from 0.003 to 0.056 g·g⁻¹·d⁻¹.

RGR was significantly ($P < 0.05$, t test) greater in subplots with neighbors removed than in subplots with neighbors left intact for 30 of the 44 plots (Fig. 1). At three sites (Guelph, Canberra, Umeå), removing neighbors increased RGR significantly in all plots. At other sites, removing neighbors increased RGR significantly only in plots with the most neighbor biomass (Morgantown, Groningen) or with the least neighbor biomass (Almonte, Vancouver) or with a medium to high amount of neighbor biomass (Long Island, Sawyerville).

Competition intensity

For all sites combined, the relationship between CI and neighbor biomass was not the same for CI_a (Fig. 2A) as for CI_c (Fig. 2B). CI_a increased significantly in a nonlinear fashion as neighbor biomass increased (Table 3). In contrast, CI_c did not change significantly as neighbor biomass increased.

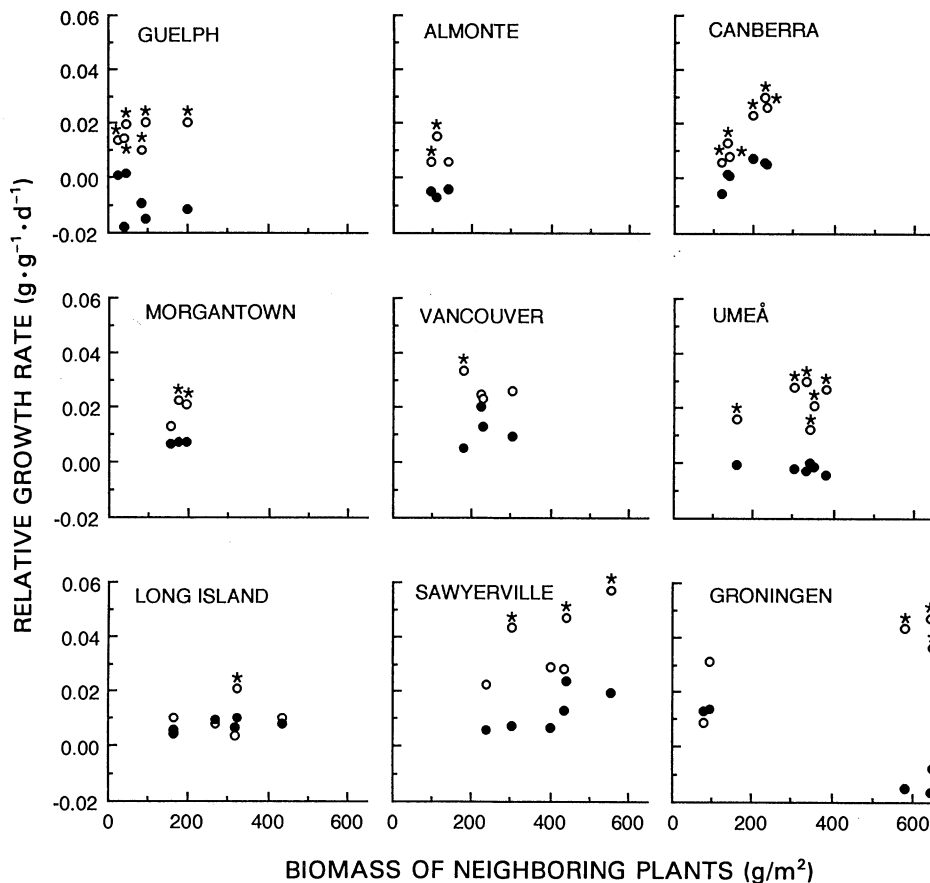


FIG. 1. Relationship between neighbor biomass and mean relative growth rate (RGR) for *Poa* transplants in subplots with neighbors removed (○) or left intact (●) at each of nine sites. * $P < 0.05$ for t test comparing RGR values for subplots with neighbors removed vs. neighbors left intact at a particular value of neighbor biomass.

When sites were considered individually, CI_a increased significantly with increasing neighbor biomass only at Groningen ($P = 0.025$, $R^2 = 0.85$, linear regression). Similarly, CI increased significantly with increasing neighbor biomass only at Groningen ($P = 0.048$, $R^2 = 0.78$, linear regression).

DISCUSSION

Our results show that competition from neighbors may reduce plant growth over a wide range of site productivity, including sites with even very low productivity. However, our results are only partly consistent with the null hypothesis that CI is independent of neighbor biomass. CI for *Poa* was independent of neighbor biomass over the relatively narrow range of values found at the majority of sites studied here, plus at two sites studied previously (DiTommaso and Aarssen 1991, Wilson and Tilman 1991). In contrast, CI was directly related to neighbor biomass at Groningen, where the range of neighbor biomass was greater than at any other site. Our finding that the relationship between CI and neighbor biomass may depend on the range of neighbor biomass considered is important for

two reasons. First, it helps to explain why previous neighbor removal experiments, each involving a different range of habitat productivity, did not all detect the same relationship between CI and neighbor biomass. Second, it confirms that neighbor biomass is more likely to be a useful predictor of variation in CI over a wide range of habitat productivity than a narrow range. Moore and Keddy (1989) made a similar point about the utility of plant standing crop as a predictor of species richness. A predicted curvilinear relationship between species richness and plant standing crop was only evident over a wide range of plant standing crop, spanning different types of communities. Species richness and standing crop were unrelated within each community type.

Our inability to reject the null hypothesis for most sites may also reflect the relatively low number of degrees of freedom (i.e., 2–5) available for the statistical test at individual sites. Either a relatively large sample size or a large range of variation in neighbor biomass, or both, may be needed to detect any effect of neighbor biomass on CI .

Theories that predict that CI is unrelated to habitat

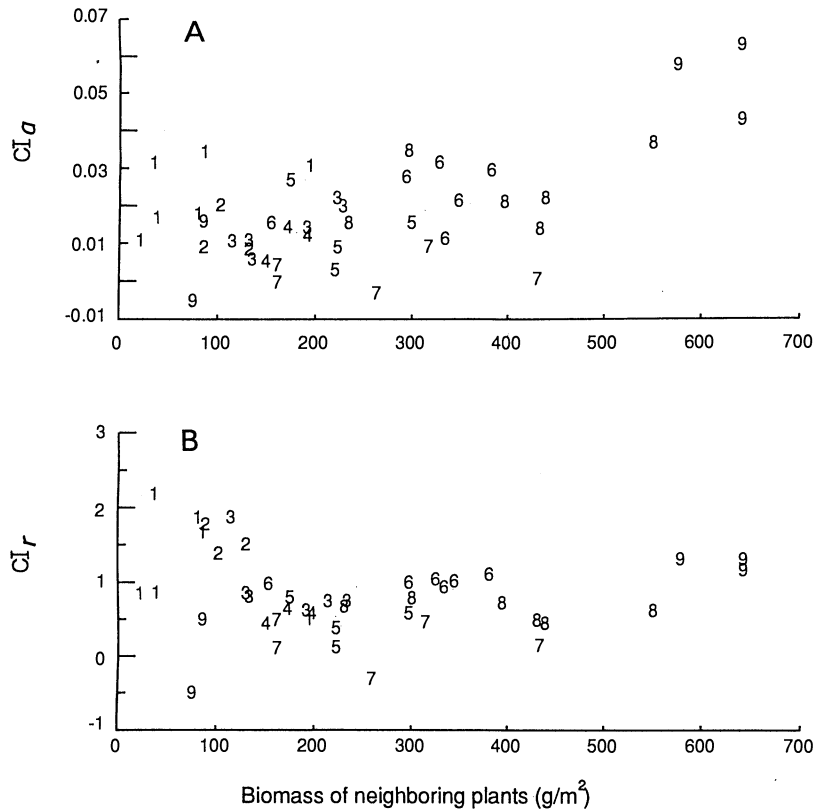


FIG. 2. Relationship between (A) neighbor biomass and absolute competition intensity (CI_a) and (B) neighbor biomass and relative competition intensity (CI_r) experienced by *Poa* transplants in 44 plots. Numbers 1 through 9 refer to site locations given in Table 1.

productivity (Tilman 1988, Taylor et al. 1990) are consistent with *Poa*'s response to neighbor removal at sites with a narrow range of neighbor biomass but not at Groningen where there was a wider range of neighbor biomass. The reverse is true for theories that predict CI is directly related to habitat productivity (Grime 1979, Keddy 1990).

Attempts to test predicted relationships between CI and habitat productivity have been complicated by imprecise definition of CI and, consequently, by dis-

agreement about how CI should be measured and calculated (Welden and Slauson 1986, Campbell et al. 1991, Grace 1991). We measured CI by removing neighbors of a target plant since our objective was to help clarify inconsistent results of previous neighbor removal experiments. Campbell et al. (1991) questioned the use of removal experiments to study competition because nutrient supply in removal subplots may increase from decomposition of dead neighbors rather than just from reduced uptake of nutrients by neighbors. Both mechanisms reflect positive effects of neighbor removal on nutrient supply so this criticism is not entirely justified (cf. Aarssen and Epp 1990). At present, we do not know the relative importance of root decomposition vs. reduced nutrient uptake by neighbors for *Poa*'s response to removal of its neighbors. The amount of nutrient released from decomposing roots would depend on their element content (e.g., N vs. P) and on abiotic conditions (Seastedt 1988, Eason and Newman 1990).

We calculated CI using two different indices and the choice of index affected the interpretation of results for all sites combined; CI and neighbor biomass were significantly related when CI was expressed in absolute terms (CI_a) but not in relative terms (CI_r). Campbell and Grime (1992) also found that CI_a was more sen-

TABLE 3. Generalized additive model (GAM) results for all sites combined, with neighbor biomass (NB) and/or site (S) as independent variables and either absolute competition intensity (CI_a) or relative competition intensity (CI_r) as the dependent variable.

De- pend- ent vari- able	Independ- ent variables	df	D^*	R^2	P values for independent variables	
					S	NB
CI_a	NB + S	29	2524.	0.70	<0.0001	0.0003
	S	34	5230.	0.37	<0.002	
CI_r	NB + S	29	3.6	0.73	<0.0001	0.35
	S	34	5.6	0.58	<0.0001	

* D = deviance, R^2 = fraction of variation explained.

sitive than CI, to a change in neighbor biomass. In both their study and ours, the absolute difference in plant mass (or RGR) with neighbors present vs. without neighbors present increased more rapidly than the relative difference. Since the choice of CI index may affect interpretation of results and different indices have been used in past studies, it is not surprising that study results have been inconsistent. Clarifying theoretical predictions about expected changes in both absolute and relative CI with habitat productivity may be useful.

In summary, our multisite study has demonstrated that competition from neighbors may influence plant growth over a wide range of site productivity and in this case there appeared to be no general relationship between CI and neighbor biomass.

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